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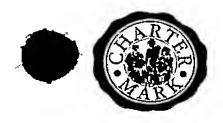
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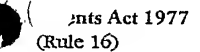
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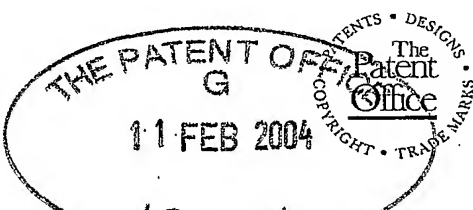
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2. Patent application number (The Patent Office will fill this part in)

0403041.7

9 9 FEB 2004

3. Full name, address and postcode of the or of each applicant (underline all surnames)

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Patents ADP number (if you know it)

If the applicant is a corporate body, give the country/state of its incorporation

8189383002

4. Title of the invention

INDUCTION OF APOPTOSIS

5. Name of your agent (if you have one)

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#### Induction of apoptosis

#### Field of the Invention

This invention relates to the induction of apoptosis by inhibition of SIRT1 expression, in particular the induction of apoptosis in tumour cells. Materials and methods for inhibiting SIRT1 expression are provided, including RNA interference methods.

#### Background to the Invention

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Many different types of stress stimulate cellular signalling pathways that result in stabilisation and activation of the tumour suppressor p53 (reviewed in Pluquet and Hainaut, 2001). Stabilisation of p53 is invariably accompanied by extensive post-translational modifications, including phosphorylation and acetylation (see Appella and Anderson, 2001). Mapping the precise relationships between stress stimuli, specific modifications and the stabilisation and activation of p53 has proven extremely difficult; however general patterns and more specific correlations are now established, particularly with regard to phosphorylation of the amino-terminus and acetylation of the carboxy-terminus.

The transcriptional activity of p53, with particular regard to its pro-apoptotic functions, is tightly regulated. Therefore if acetylation functions to activate p53 transcriptional activity, a logical assumption would be that p53 acetylation is subject to negative control.

Until recently, deacetylation of p53 was only known to be performed by members of the trichostatin A-sensitive histone

deacetylase (HDAC) class I family (Juan et al., 2000; Luo et al., 2000). Indeed, mounting evidence suggests that p53 utilises these HDACs to repress specific promoters (Murphy et al., 1999). More recently, the human sirtuin SIRT1 (Frye, 1999) has been identified as a bona fide p53 deacetylase (Luo et al., 2001; Vaziri et al., 2001).

The sirtuins are a ubiquitous gene family found throughout eukarya and prokarya, defined by conserved ~250 amino acid core domain. Many of the sirtuins are NAD-dependent deacetylases ['NDAC'].

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The function of Saccharomyces cerevisiae sirtuin SIR2 has been extensively studied. It has many activities including silencing of mating-type loci, telomeric position effect silencing and silencing at the rDNA locus, suppression of illegitimate recombination and increasing cellular control of longevity. It is also implicated in response to dsDNA breaks.

Humans have seven sirtuins, although not all appear to have NDAC activity.

Human SIRT2 is a cytoplasmic, microtubule-associated protein.

Is shows increases in abundance and phosphorylation at G2/M. It

is a tubulin deacetylase and is strongly down-regulated in many
gliomas and glioma cell lines. Transgene replacement causes
microtubule disruption and strongly reduces the number of stable
clones expressing SIRT2 compared to a control in colony
formation assays )Hiratsuka, M et al (2003) Biochem Biophys Res

Commun. 309(3) 558-566).

Human SIRT3 is synthesized as an inactive proenzyme and activated by proteolysis on insertion into the mitochondrial matrix. Its function is unknown

- 5 Human SIRT1 is the closest human homologue to yeast SIR2. It is a nuclear protein found throughout the nucleus. Immunostaining of cells with anti-SIRT1 antibodies shows diffuse nuclear staining.
- SIRT1 interacts with p53 via the p53 core and carboxy-terminus. It appears to act as a p53 deacetylase, as overexpression of SIRT1 results in reduced acetylation of p53. This in turn leads to reduced expression of endogenous p21, reduced transcription from a p21 reporter construct, and reduced apoptosis in response to  $\rm H_2O_2$  and  $\rm \gamma\text{-}rays$ .

Overexpression of a catalytically inactive mutant SIRT1 enhances acetylation of p53, and sensitises cells to apoptosis induced by  $\rm H_2O_2$  and  $\rm \gamma\text{-}rays$ .

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The current consensus is that SIRT1 negatively regulates p53 function via deacetylation of p53, so that inhibition of SIRT1 function sensitises cells to p53-dependent apoptosis in response to cellular stress (Luo et al., 2001; Vaziri et al., 2001;

25 Langley et al, 2002)

However, modulation of SIRT1 activity has until now been achieved by treatment of cells with the SIRT1 inhibitor nicotinamide, or by overexpression in trans of wild type and catalytically inactive forms of SIRT1. The use of nicotinamide is problematic, partly due to potential inhibition of the SIRT1-related NDACs and SIRT2 and SIRT3, and partly due to potential

pleiotropic effects as nicotinamide is a natural cellular intermediary metabolite. Recent studies have also shown that transgene dosage may be critical in the analysis of p53 function (Blattner et al., 1999; Dumaz et al., 2001), and may underlie the sometimes-conflicting results that have been reported for p53 using this technique.

#### Summary of the Invention

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In order to elucidate the role of SIRT1 in regulating p53

deacetylation, p53 function and apoptosis, we sought a specific and efficient method of inhibiting the expression or activity of SIRT1.

We addressed the problems of the prior art methods by using a 15 more specific technique, RNA interference, to inhibit SIRT1 expression. RNA interference (RNAi) was used to analyse the function of p53 carboxy-terminal acetylation in p53-dependent apoptosis. The aim was to stabilise acetylation of p53 by silencing of SIRT1. RNAi is a sequence-specific posttranslational gene silencing mechanism, which can be initiated - 20 in cultured mammalian cells by transfection of a 19-21 nucleotide RNA duplex (short interfering RNA; 'siRNA') homologous in sequence to the target mRNA (Elbashir et al., Importantly, it has been demonstrated that RNAi does not 25 in itself engage the apoptotic machinery nor alter apoptotic processes, (Jiang and Milner, 2002; Jiang and Milner 2003).

Using RNAi to inhibit SIRT1 expression, we surprisingly found that inhibition of SIRT1 induces massive apoptosis in tumour cells even without additional apoptotic stimuli. This effect is independent of p53 and thus appears to represent a completely

new and unexpected activity of SIRT1. It also appears to be independent of the known pro-apoptotic proteins Bax and PUMA.

Even more surprisingly, the induction of apoptosis seen on inhibition of SIRT1 expression by RNAi appears to be specific to tumour cells, as it does not occur in normal human fibroblasts. This opens the way for new approaches to cancer treatment via the specific induction of apoptosis in tumour cells.

10 Prior to the invention, inhibition of SIRT1 expression would not have been expected to induce apoptosis in tumour cells in preference to normal cells. In fact, the opposite assumption would have been made, as it was thought that SIRT1 acted only on the p53-dependent pathway of apoptosis, and many tumour cells lack functional p53. Further, it would not have been expected 15 that inhibition of SIRT1 expression alone would induce apoptosis in the absence of other stimuli to trigger p53-dependent apoptotic pathways. The observations of the prior art indicated that inhibition of SIRT1 expression would exacerbate the 20 apoptotic phenotype of cells dying in response to agents that activate p53-dependent apoptotic pathways (Luo et al., 2001; Vaziri et al., 2001), but not that loss of SIRT1 expression itself could induce apoptosis in a p53-independent manner

The invention accordingly provides for a method of inducing apoptosis in a cell comprising administering a SIRT inhibitor to the cell. Preferably, the cell is a tumour cell. In some embodiments, the cell lacks functional p53, Bax and/or PUMA protein.

The SIRT1 inhibitor may be an agent for inducing RNA interference in a cell, such as a siRNA, a dsRNA, or a nucleic

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acid encoding such RNA. In a preferred embodiment, said agent is a siRNA.

In another aspect, the invention provides a method of treating a proliferative disease comprising administering to an individual in need thereof an effective amount of a SIRT1 inhibitor. The disease may be cancer, for example a colorectal carcinoma.

Thus, the invention also provides a SIRT1 inhibitor for use in a method of medical treatment or therapy. The therapy may be treatment of a proliferative disease, for example cancer. Also provided is the use of a SIRT1 inhibitor in the manufacture of a medicament for the treatment of a proliferative disease.

- In another aspect, the invention provides a method of identifying a SIRT inhibitor for use in a method of treatment of the invention, the method comprising administering a candidate compound to cultured tumour cells in vitro; determining whether SIRT expression and/or activity is reduced in said cells; and assaying for apoptosis of said cells. The method may further comprise a step of administering said candidate compound to cultured non-tumour cells in vitro and assaying for apoptosis of said cells.
- In another aspect, the invention provides an agent for inhibiting the expression of SIRT1 protein in a cell. The agent may be an agent which induces RNA interference to SIRT1 mRNA in a cell, for example siRNA, a dsRNA, or a nucleic acid encoding such RNA. In a preferred embodiment, the agent is a siRNA.

Compositions for pharmaceutical use comprising agents of the invention in combination with a pharmaceutically acceptable excipient are also provided.

5 The invention will now be described in more detail with reference to the following figures.

### Brief description of the Figures

Figure 1. siRNAs used in this study.

SIRT1 siRNA (SEQ ID No. 1)

Lamin A/C siRNA

BCR-Abl siRNA

#### Figure 2

- 15 A. RT-PCR showing that SIRT siRNA suppresses SIRT-1 mRNA expression.
  - B. RT-PCR of SIRT-1 in p53<sup>+/+</sup> cells (top) and p53<sup>-/-</sup> cells (bottom) transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated.
  - C. Control: RT-PCR of vimentin in p53\*/\* cells (left) and p53\*/\* cells (right) transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated.
    - D. Control: RT-PCR of Lamin A/C in p53\*/\* cells (left) and p53\*/- cells (right) transfected with SIRT1 siRNA, Lamin A/C siRNA
  - 25 and Bcr-Abl siRNA as indicated.
    - Figure 3. Western blots showing that SIRT1 siRNA suppresses SIRT1 protein expression.
  - A. Western blot of whole cell lysates from cells transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as

indicated. Blots were probed with an anti-SIRT1 antibody and ananti-Lamin A/C antibody as shown.

B. Western blot of the soluble fraction of lysates from p53<sup>+/+</sup> cells (left) and p53<sup>-/-</sup> cells (right) transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated. Equal amounts of total protein were loaded in each lane. Blots were probed with an anti-SIRT1 antibody. The bands 2' and 10' indicate 2 minute and 10 minute exposure times respectively C. Western blot of the soluble fraction of lysates from p53<sup>+/+</sup> cells transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated. The amount of total protein in each lane was not equalised. Blots were probed with an anti-SIRT1 antibody.

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- D. Western blot of the soluble fraction of lysates from p53-/cells transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated. The amount of total protein in each lane was not equalised. Blots were probed with an anti-SIRT1 antibody.
- Figure 4 Western blots showing that SIRT1 siRNA stabilises p53 protein and increases p53 protein expression.
  - A. Western blot of whole cell lysates, soluble and insoluble fractions from cells transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated. Blots were probed with an anti-p53 antibody.
- B. Western blot of whole cell lysates (WCE), soluble (SOL) and insoluble (PEL) fractions of lysates from p53<sup>+/+</sup> cells (top) and p53<sup>-/-</sup> cells (bottom) transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated. Blots were probed with an anti-p53 antibody. Recombinant human p53 was run as an antibody control.

- C. Western blot of whole cell lysates (WCE), soluble (SOL) and insoluble (PEL) fractions of lysates from p53<sup>+/+</sup> cells transfected with SIRT1 siRNA (top), Lamin A/C siRNA (middle) and Bcr-Abl siRNA (bottom). Blots were probed with an anti-p53 antibody. Recombinant human p53 was run as an antibody control.
- Figure 5 Western blots showing that SIRT1 siRNA induces phosphorylation of p53 protein at serine 15.
- A. Western blots of whole cell lysates, soluble and insoluble fractions from cells transfected with SIRT1 siRNA. Blots were probed with the anti-p53 antibody DO-1 (top) and an anti-phosphoSer15 antibody (middle). The bottom blot was probed with antibodies to p21 and HDM2. Cells were harvested at 12, 24, 36 and 48 hours post-transfection, as indicated.
- 15 B. Western blot of whole cell lysates (WCE), soluble (SOL) and insoluble (PEL) fractions of lysates from p53<sup>+/+</sup> cells transfected with SIRT1 siRNA (top right), Lamin A/C siRNA (bottom left) and Bcr-Abl siRNA(bottom right). Blots were probed with an anti-phosphoSer15 antibody. Recombinant human p53 was run as an antibody control.
  - C. Western blot of whole cell lysates (WCE) from cells transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated. Blots were probed with an anti-phosphoSer15 antibody.

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- Figure 6. Western blots showing that SIRT1 siRNA induces expression of p21.
- A. Western blot of soluble lysates from p53<sup>+/+</sup> cells (left) and p53<sup>-/-</sup> cells (right) transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated. Blots were probed with an anti-p21 antibody.

- B. Western blot of whole cell lysates from p53<sup>+/+</sup> cells incubated at 37°C (bottom) and subjected to cold shock (top) and transfected with SIRT1 siRNA, Lamin A/C siRNA and Bcr-Abl siRNA as indicated. Blots were probed with an anti-p21 antibody.
- 5 SIRT1 inhibition does not induce p21 expression under conditions of cold shock.

Figure 7 Controls: Effect of SIRT siRNA on expression of Lamin A/C and Bcr-Abl.

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- Figure 8 SIRT siRNA induces apoptosis in p53<sup>+/+</sup> tumour cells.
- A Phase contrast images of HCT116 p53\*/\* cells 24 hours and 48 hours after transfection with SIRT1 siRNA, BCR-Abl siRNA and Lamin A/C siRNA.
- 15  $\bf B$  Phase contrast images of HCT116 p53<sup>+/+</sup> cells at 12 hour timepoints after transfection with SIRT siRNA.
  - Figure 9 SIRT siRNA induces apoptosis in p53+/+ tumour cells.
- A Phase contrast images of HCT116 p53-/- cells 24 hours, 48 hours after transfection with SIRT1 siRNA, BCR-Abl siRNA and Lamin A/C siRNA.
- Figure 10 SIRT siRNA induces apoptosis in Bax<sup>-/-</sup> tumour cells.

  A Phase contrast images of HCT116 p53<sup>+/+</sup> Bax<sup>-/-</sup> cells 24 hours

  and 48 hours and after transfection with SIRT1 siRNA.
  - Figure 11 FACS analysis showing apoptosis induced in SIRT1transfected HCT116 p53<sup>+/+</sup>·cells. Cells were analysed at 48 hours post-transfection.

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Figure 12 Western blots showing Bax and PUMA expression in cells treated with SIRT1 siRNA. Blots of lysates from cells

transfected with SIRT1 siRNA were probed with an anti-Bax antibody and an anti-PUMA antibody as shown.

Figure 13 SIRT1 siRNA does not induce apoptosis in primary human fibroblasts.

Phase contrast images of primary human normal diploid fibroblasts (NDFs) transfected with SIRT1 siRNA, taken at 2, 3, 4 and 5 days post-transfection.

10 Figure 14 cDNA sequence of human SIRT1 (SEQ ID No. 2).

Figure 15 Amino acid sequence alignment of human SIRT1 (SEQ ID No. 3), 2 and 3. Asterisks indicate conserved residues. ':' represents conservative amino acid substitution and '.' represents semi-conservative amino acid replacement.

Figure 16 cDNA sequence alignment of human SIRT1-7 to either side of the region of SIRT 1 covered by the SIRT-1 siRNA (shown in bold) to demonstrate specificity of SIRT1 siRNA sequences..

Figure 17 Amino acid sequence alignment of human SIRT1-7.

#### Detailed description

#### Proteins

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25 SIRT1 refers to human sirtuin 1 as described above. SIRT1 is encoded by a nucleic acid with nucleotide sequence shown in SEQ ID No. 2 and Fig. 14. The full-length SIRT1 protein has the sequence shown in SEQ ID No. 3 and Fig. 15. The GenBank accession number for the full-length SIRT1 cDNA and amino acids sequences is NM012238.

Functional p53 protein indicates p53 protein which is capable of inducing apoptosis. A cell lacking functional p53 protein may, for example, carry a deletion or termination mutation of p53 so that it does not express full length p53 at all. Alternatively, the cell may express a mutant p53 which is not capable of triggering apoptosis.

PUMA (p53 upregulated modulator of apoptosis) is a mitochondrial protein that appears to be required for p53-mediated apoptosis (Yu et al., 2003). Expression of PUMA is tightly regulated by p53. PUMA triggers apoptosis via Bax by binding to Bcl-2 and Bcl-X<sub>L</sub>, which otherwise bind and sequester Bax.

Bax is member of the Bcl-2 family of apoptosis regulatory

15 proteins which induces apoptosis, at least in part by triggering release of cytochrome c from mitochondria. Cytochrome c mediates the subsequent activation of the caspases which carry out the apoptotic death programme. Expression of Bax may also be upregulated by p53.

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#### SIRT1 inhibitor

The term 'SIRT1 inhibitor' is intended to cover any agent that reduces the expression or activity of SIRT1 in a cell.

Alternatively, the agent may be an agent that inhibits the transcription or translation of SIRT, such as an antisense DNA, RNA or an agent that induces RNA interference.

Inhibition of SIRT expression may be detected by RT-PCR using SIRT1-specific primers, or by Western blotting using an anti-SIRT1 antibody

#### RNA interference

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RNA interference (RNAi) is a process whereby the introduction of double stranded RNA (dsRNA) into a cell inhibits gene expression post-transcriptionally, in a sequence dependent fashion. This process is also known as post-transcriptional gene silencing. Current models of RNAi indicate that it is mediated by short (typically 20-25 nucleotides) dsRNAs known as 'small interfering RNAs' (siRNA). It appears that dsRNA is cleaved in the cell to create siRNAs. siRNAs are then incorporated into an RNA-induced 10 silencing complex (RISC), guiding the complex to the homologous endogenous mRNA. The activated RISC then cleaves the mRNA transcript, resulting in the destruction of the mRNA in a cell which is homologous to the siRNAs. The siRNAs are re-cycled. In this way, a relatively small number of siRNAs can selectively 15 destroy a large excess of cellular mRNA.

To induce RNA interference in a cell, dsRNA may be introduced into the cell as an isolated nucleic acid fragment or via a transgene, plasmid or virus. Alternatively, siRNA may be synthesised and introduced directly into the cell.

siRNA sequences are selected on the basis of their homology to the gene it is desired to silence. Homology between two nucleotide sequences may be determined using a variety of programs including the BLAST program, of Altschul et al. (1990) J. Mol. Biol. 215: 403-10, or BestFit, which is part of the Wisconsin Package, Version 8, September 1994, (Genetics Computer Group, 575 Science Drive, Madison, Wisconsin, USA, Wisconsin 53711). Sequence comparisons may be made using FASTA and FASTP (see Pearson & Lipman, 1988. Methods in Enzymology 183: 63-98). Parameters are preferably set, using the default matrix, as

follows: Gapopen (penalty for the first residue in a gap): -16 for nucleic acid; Gapext (penalty for additional residues in a gap): -4 for nucleic acids; KTUP word length: 6 for nucleic acids.

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Sequence comparison may be made over the full length of the relevant sequence, or may more preferably be over a contiguous sequence of about or 10, 15, 20, 25 or 30 bases.

Preferably the degree of homology between the siRNA and the target gene is at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, or at least 99%.

The degree of homology between the siRNA or dsRNA and the gene to be silenced will preferably be sufficient that the siRNA or dsRNA will hybridise to the nucleic acid of the gene sequence under stringent hybridisation conditions.

Typical hybridisation conditions use 4-6 x SSPE; 5-lox Denhardts solution, 5g polyvinylpyrrolidone and 5g bovine serum albumin; 100µg-lmg/ml sonicated salmon sperm DNA; 0.1-1% sodium dodecyl sulphate; optionally 40-60% deionised formamide. Hybridisation temperature will vary depending on the GC content of the nucleic acid target sequence but will typically be between 42°C-65°C.

25 Sambrook et al (2001) Molecular Cloning: A Laboratory Approach (3<sup>rd</sup> Edn, Cold Spring Harbor Laboratory Press). A common formula for calculating the stringency conditions required to achieve hybridisation between nucleic acid molecules of a specified homology is:

30  $T_m = 81.5$ °C+16.6Log[Na<sup>+</sup>]+0.41[%G+C]-0.63(% formamide).

The siRNA may be between 10bp and 30bp in length, preferably

between 20bp and 25bp. Preferably, the siRNA is 19, 20, 21 or 22bp in length.

The siRNA sequence may be, for example, any suitable contiguous sequence of 10-30bp from the sequence shown in Fig 14 (SEQ ID No. 2). Alternatively, longer dsRNA fragments comprising contiguous sequences from the sequences of Fig 14 (SEQ ID No. 2) may be used, as they will be cleaved to form siRNAs within the cell. Preferably, the siRNA sequence is that shown in Fig 1a (SEQ ID No. 1).

In a preferred embodiment, the siRNA has the 19bp sequence shown in Fig 1. In some embodiments, the siRNA has an overhang at one or both ends of one or more deoxythymidine bases. The overhang is not to be interpreted as part of the siRNA sequence. Where present, it serves to increase the stability of the siRNA within cells by reducing its susceptibility to degradation by nucleases.

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siRNA molecules may be synthesized using standard solid or solution phase synthesis techniques which are known in the art. Linkages between nucleotides may be phosphodiester bonds or alternatives, for example, linking groups of the formula P(O)S, (thioate); P(S)S, (dithioate); P(O)NR'2; P(O)R'; P(O)OR6; CO; or CONR'2 wherein R is H (or a salt) or alkyl (1-12C) and R6 is alkyl (1-9C) is joined to adjacent nucleotides through-O-or-S-.

Alternatively, siRNA molecules or longer dsRNA molecules may be made recombinantly by transcription of a nucleic acid sequence, preferably contained within a vector as described below.

Modified nucleotide bases can be used in addition to the

naturally occurring bases, and may confer advantageous properties on siRNA molecules containing them.

For example, modified bases may increase the stability of the siRNA molecule, thereby reducing the amount required for silencing. The provision of modified bases may also provide siRNA molecules which are more, or less, stable than unmodified siRNA.

The term 'modified nucleotide base' encompasses nucleotides with a covalently modified base and/or sugar. For example, modified nucleotides include nucleotides having sugars which are covalently attached to low molecular weight organic groups other than a hydroxyl group at the 3'position and other than a phosphate group at the 5'position. Thus modified nucleotides may also include 2'substituted sugars such as 2'-O-methyl-; 2-O-alkyl; 2'-S-alkyl; 2'-S-allyl; 2'-fluoro-; 2'-halo or 2; azido-ribose, carbocyclic sugar analogues a-anomeric sugars; epimeric sugars such as arabinose, xyloses or lyxoses, pyranose sugars, furanose sugars, and sedoheptulose.

Modified nucleotides are known in the art and include alkylated purines and pyrimidines, acylated purines and pyrimidines, and other heterocycles. These classes of pyrimidines and purines are known in the art and include pseudoisocytosine, N4,N4-ethanocytosine, 8-hydroxy-N6-methyladenine, 4-acetylcytosine,5-(carboxyhydroxylmethyl) uracil, 5 fluorouracil, 5-bromouracil, 5-carboxymethylaminomethyl-2-thiouracil, 5-carboxymethylaminomethyl uracil, dihydrouracil, inosine, N6-isopentyl-adenine, 1- methyladenine, 1-methylpseudouracil, 1-methylguanine, 2,2-dimethylguanine, 2methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-

methyladenine, 7-methylguanine, 5-methylaminomethyl uracil, 5methoxy amino methyl-2-thiouracil, -D-mannosylqueosine, 5methoxycarbonylmethyluracil, 5methoxyuracil, 2 methylthio-N6isopentenyladenine, uracil-5-oxyacetic acid methyl ester,
5 psueouracil, 2-thiocytosine, 5-methyl-2 thiouracil, 2thiouracil, 4-thiouracil, 5methyluracil, N-uracil-5-oxyacetic
acid methylester, uracil 5-oxyacetic acid, queosine, 2thiocytosine, 5-propyluracil, 5-propylcytosine, 5-ethyluracil,
5ethylcytosine, 5-butyluracil, 5-pentyluracil, 5-pentylcytosine,
and 2,6,diaminopurine, methylpsuedouracil, 1-methylguanine, 1methylcytosine.

#### Vectors

30

The invention also provides vectors comprising a nucleotide

15 sequence encoding an siRNA or longer RNA or DNA sequence for production of dsRNA. The vector may be any RNA or DNA vector. The vector is preferably an expression vector, wherein the nucleotide sequence is operably linked to a promoter compatible with the cell. The vector will preferably have at least two

20 promoters, one to direct expression of the sense strand and one to direct expression of the antisense strand of the dsRNA. Alternatively, two vectors may be used, one for the sense strand and one for the antisense strand. Alternatively the vector may encode RNAs which form stem-loop structures which are

25 subsequently cleaved by the cell to produce dsRNA.

Where the vector is an expression vector, the sequence to be expressed will preferably be operably linked to a promoter functional in the target cells. Promoters suitable for use in various vertebrate systems are well known. For example, suitable promoters include viral promoters such as mammalian

retrovirus or DNA virus promoters, e.g. MLV, CMV, RSV, SV40 IEP and adenovirus promoters and metallothionein promoter. The CMV IEP may be more preferable for human use. Strong mammalian promoters may also be suitable as well as RNA polymerase II and III promoters. Variants of such promoters retaining substantially similar transcriptional activities may also be used.

Other vehicles suitable for use in delivering nucleic acids such as siRNAs include viruses and virus-like particles (VLPs) such as HPV VLPs comprising the L1 and/or L2 HPV viral protein; or hepatitis B viral proteins. Other suitable VLPs may be derived from picornaviruses; togaviruses; rhabdoviruses; orthomyxoviruses; retroviruses; hepadnaviruses; papovaviruses; adenoviruses; herpesviruses; and pox viruses.

#### Delivery

Various agents may be used to improve the delivery of RNA, DNA or protein into the cell. Viral vectors as described above may be used to deliver nucleic acid into a cell. Where other vectors, or no vector, is used, delivery agents such as liposomes may usefully be employed. Delivery peptides such as Antennapedia of the HIV TAT peptide may be used, as may organic polymers such as a dendrimers or polylysine-transferrine
25 conjugates.

Liposomes can be prepared from a variety of cationic lipids, including DOTAP, DOTMA, DDAB, L-PE, and the like. Lipid carrier mixtures containing a cationic lipid, such as N-[1-(2,3-dioleyloxy) propyl]-N,N,N-triethylammonium chloride (DOTMA) also known as "lipofectin", dimethyl dioctadecyl ammonium bromide

(DDAB), 1,2-dioleoyloxy-3-(trimethylammonio) propane (DOTAP) or L-lysinyl-phosphatidylethanolamine (L-PE) and a second lipid, such as dioleoylphosphatidylethanolamine (DOPE) or cholesterol (Chol), are particularly useful for use with nucleic acids.
DOTMA synthesis is described in Felgner, et al., (1987) Proc. Nat. Acad. Sciences, (USA) 84:7413-7417. DOTAP synthesis is described in Stamatatos, et al., Biochemistry, (1988) 27:3917-3925.

Liposomes are commercially available from many sources.

DOTMA:DOPE lipid carriers can be purchased from, for example,
BRL. DOTAP:DOPE lipid carriers can be purchased from Boehringer
Mannheim. Cholesterol and DDAB are commercially available from
Sigma Corporation. DOPE is commercially available from Avanti
Polar Lipids. DDAB:DOPE can be purchased from Promega.

Invitrogen make liposomes under the names oligofectamine™ and
lipofectamine™.

To incorporate nucleic acid into liposomes, the liposome-nucleic acid complex is prepared by mixing with the nucleic acid in an appropriate nucleic acid:lipid ratio (for example 5:3) in a physiologically acceptable diluent (for example Opti-MEM<sup>TM</sup> at an appropriate dilution) immediately prior to use.

#### 25 Apoptosis assays

The induction of apoptosis in cells may be assayed by many methods.

Apoptotic cells in culture may be detected and assayed by
30 photomicrography; apoptotic cells may be detected by their
distinctive morphology, with blebbing of the plasma membrane and

chromatin condensation and fragmentation. DNA dyes such as propidium iodide (PI) or Hoechst 33342 may be used to detect chromatin condensation. Alternatively, TUNEL may be used to detect DNA strand breaks.

5

Antibodies such as anti-annexin V can be used to label apoptotic cells and detected by immunofluorescence, and assayed by micrography of FACS analysis.

10 FACS analysis may be used in combination with a DNA dye such as PI, TUNEL and/or annexin staining. Apoptotic cells may be detected as a sub-G1 fraction of cells which have lost DNA.

Release of cytochrome c from mitochondria is another marker for apoptosis. A measure of the early stages of apoptosis may be obtained by fractionating cell lysates into mitochondrial and cytoplasmic fractions and detecting the amount of cytochrome c released from the mitochondria into the cytosol. This is usually done by performing a Western blot and probing with an anti-cytochrome c antibody.

#### Proliferative disease

A proliferative disease is a pathological condition characterised by unwanted cell growth. In general,

25 proliferative diseases can be divided into two types: clonal and non-clonal. Clonal proliferative disease usually leads to the formation of tumours, which may be benign or malignant (cancerous). Cancers may be cancer of the skin, connective tissue, adipose, breast, lung, stomach, pancreas, ovary, cervix, uterus, kidney, bladder, colon, prostate, central nervous system (CNS), retina and circulating tumours (such as leukaemia and

lymphoma). Colorectal cancer includes cancers of the colon, rectum, anus, and appendix.

SIRT inhibitors may be effective in providing treatments that discriminate between malignant and normal cells, avoiding many of the deleterious side-effects present with most current chemotherapeutic regimes.

Non-clonal proliferative diseases include psoriasis, fibrocystic disease, myelofibrosis, proliferative diabetic retinopathy, atherosclerosis (associated with proliferation of vascular cells) and chronic inflammatory proliferative diseases (CIPD).

As used herein, 'tumour cells' shall be taken to refer both to cells derived from tumours, including malignant tumours, and cells immortalised in vitro. 'Normal' cells refers to cells with normal growth characteristics that do not show abnormal proliferation.

'Therapy' and 'treatment' of disease includes any therapy or treatment that alleviates in any way the symptoms of a disease. These terms refer to any administration of the compound, salt or N-oxide thereof, intended to alleviate the severity of a disorder of the GI tract in a subject, and includes treatment intended to cure the disease, provide relief from the symptoms of the disease and to prevent or arrest the development of the disease in an individual at risk from developing the disease or an individual having symptoms indicating the development of the disease in that individual.

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#### Compositions

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Pharmaceutical compositions according to the present invention, and for use in accordance with the present invention, may include, in addition to active ingredient, a pharmaceutically acceptable excipient, carrier, buffer, stabiliser or other materials well known to those skilled in the art. Such materials should be non-toxic and should not interfere with the efficacy of the active ingredient. The precise nature of the carrier or other material will depend on the route of administration, which may be oral, or by injection, e.g. cutaneous, subcutaneous or intravenous.

For solid compositions, conventional non-toxic solid carriers include, for example, pharmaceutical grades of mannitol, lactose, cellulose, cellulose derivatives, starch, magnesium 15 stearate, sodium saccharin, talcum, glucose, sucrose, magnesium carbonate, and the like may be used. The active compound as defined above may be formulated as suppositories using, for example, polyalkylene glycols, acetylated triglycerides and the like, as the carrier. Liquid pharmaceutically administrable 20 compositions can, for example, be prepared by dissolving, dispersing, etc, an active compound as defined above and optional pharmaceutical adjuvants in a carrier, such as, for example, water, saline aqueous dextrose, glycerol, ethanol, and the like, to thereby form a solution or suspension. If desired, 25 the pharmaceutical composition to be administered may also contain minor amounts of non-toxic auxiliary substances such as wetting or emulsifying agents, pH buffering agents and the like, for example, sodium acetate, sorbitan monolaurate, triethanolamine sodium acetate, sorbitan monolaurate, 30 triethanolamine oleate, etc. Actual methods of preparing such

dosage forms are known, or will be apparent, to those skilled in this art; for example, see Remington's Pharmaceutical Sciences, 20th Edition, 2000, pub. Lippincott, Williams & Wilkins. The composition or formulation to be administered will, in any event, contain a quantity of the active compound(s) in an amount effective to alleviate the symptoms of the subject being treated.

#### Administration

Where a composition as described herein is to be administered to an individual, administration is preferably in a "prophylactically effective amount" or a "therapeutically effective amount" (as the case may be, although prophylaxis may be considered therapy), this being sufficient to show benefit to the individual. The actual amount administered, and rate and time-course of administration, will depend on the nature and severity of what is being treated. It will also depend upon toxicity of the therapeutic agent, as determined by pre-clinical and clinical trials.

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Dosing is dependent on severity and responsiveness of the condition to be treated, with course of treatment lasting from several days to several months or until a diminution of disease state is achieved. Optimal dosing schedules are easily calculated from measurements of drug accumulation in the body. Persons of ordinary skill can easily determine optimum dosages, dosing methodologies and repetition rates. Therapeutically or prophylactically effective amounts (dosages) may vary depending on the relative potency of individual compositions, and can generally be routinely calculated based on molecular weight and EC50s in in vitro and/or animal studies. For example, given the molecular weight of an siRNA drug compound (derived from

oligonucleotide sequence and chemical structure) and an experimentally derived effective dose such as an  $IC_{50}$ , for example, a dose in mg/kg is routinely calculated. In general, dosage is from 0.001 $\mu$ g to 100g and may be administered once or several times daily, weekly, monthly or yearly.

Compositions of the present invention may be administered in a number of ways depending upon whether local or systemic treatment is desired and upon the area to be treated.

10 Administration may be topical (including ophthalmic, vaginal, rectal, intranasal, epidermal and transdermal), oral or parenteral. Parenteral administration includes intravenous drip, subcutaneous, intraperitoneal or intramuscular injection, pulmonary administration, e.g., by inhalation or insufflation, or intracranial, e.g., intrathecal or intraventricular, administration.

Prescription of treatment, e.g. decisions on dosage etc, is within the responsibility of general practitioners and other medical doctors. A composition may be administered alone or in combination with other treatments, either simultaneously or sequentially dependent upon the condition to be treated.

The invention is illustrated by the following examples.

#### Experimental procedures

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#### Design of siRNA specific to SIRT1

A siRNA sequence located adjacent to the conserved sirtuin

domain on the carboxy-terminal side was selected on the basis of

its selectivity for SIRT1 and its predicted lack of secondary

structure.

Bestfit matching to other sirtuin cDNAs was performed and the results were: SIRT2/3/7 - 57.9%

SIRT4/5/6 - 63.2%

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BLAST searching of GenBank, EMBL, DDBJ and PDB databases was performed and the results were:

Matches 1-8 : Human SIRT1 [some truncated clones].

Match 9: Streptococcus mutans sequence [imperfect].

10 Match 10: Mus musculus sequence [imperfect].

BLAST searching of the Genbank Human EST database was performed and the results were:

Matches 1-4: Human SIRT1.

15 Matches 5-10: Misc. sequences [imperfect].

These results showed that the selected sequence was likely to be specific for SIRT RNA.

The biophysical properties and predicted secondary structure formation of the selected sequence was

Tm: 46°C

GC: 42.1%

Loop: No

25 Secondary structure: No

Taken together, these results indicated that the selected sequence was likely to be effective in inducing RNA interference specific for the SIRT1 mRNA.

30

Control siRNAs were Lamin A/C siRNA (purchased from Dharmacon) and Bcr-Abl siRNA (siACE RNA, obtained from Dr Ming Jiang). A

further control, indicated in the figures as CTRL or Mock, was the transfection of siRNA buffer alone (no siRNA).

#### RT-PCR

5 RT-PCR for SIRT1, Lamin A/C, BCR-Abl and controls (vimentin and GAPDH) was performed.

A SIRT1 upstream [5'] primer located in conserved core domain, over an exon-exon boundary, and a downstream [3'] primer located in SIRT1-specific sequence were selected.

RT-PCR was performed from total RNA using a Reverse-It One-Step kit (ABgene) according to the manufacturer's instructions, and visualised on 2% TAE gels with ethidium bromide.

Primers were as follows:

SIRT1 sense 5-TCAGTGTCATGGTTCCTTTGC-3; SIRT1 antisense 5-

15 AATCTGCTCCTTTG

CCACTCT;

Lamin A sense 5-AAGCAGCGTGAGTTTGAGAGC-3;
Lamin A antisense 5-AGGGTGAACTTTGGTGGGAAC-3;
GAPDH sense 5-CGGAGTCAACGGATTTG

20 GTCGTAT-3;

25

GAPDH antisense 5-AGCCTTCTCCATGGTGGAAGAC-3.

Vimentin sense: 5-gCCAACTACATCgACAAggTg-3

Vimentin antisense: 5-qAqCAggTCTTggTATTCACg-3

Western blotting

Cells for protein lysates were washed in PBS and lysed in buffer IPAX (10 mM TRIS base pH 8.0, 140 mM NaCl, 0.5% NP40, 1 mM PMSF, 1x Complete Protease Inhibitors (Roche), 50 mM NaF, 1 mM Na<sub>3</sub>VO4, 1 mM trichostatin A, 10 mM nicotinamide). Lysates were spun at 13 000g for 20 minutes at 4°C to separate soluble and insoluble fractions, where indicated. Mitochondrial and cytosolic

fractions were prepared as previously described (Marsden et al., 2002), where indicated. Lysates were run on 10% or 15% SDS-PAGE, electroblotted to Protran membrane (Schleicher & Schuell), and probed with antibodies as described below. Visualisation was with the POD chemiluminescence kit (Roche).

Blots were probed with the following antibodies:
anti-SIRT1 (H-300, Santa Cruz); anti-p53 (D0-1, Oncogene); antiphosphoserine 15 (Ser15-R, Santa Cruz), anti-p21 (SX118,

Pharmingen); anti-HDM2 (monoclonal antibody 4B2, prepared inhouse); anti-lamin A/C (636, Santa Cruz); Anti-Abl(8E9,
Pharmingen); anti-Bax (N-20, Santa Cruz); anti-PUMAα (AHP727,
Serotec) and anti-cytochrome c (7H8.2C12, Pharmingen).

#### 15 Photomicrography

Phase contrast images were captured with an Axiovert 200M Cell Observer platform (Zeiss) at various time points posttransfection.

## 20 Cell cycle analysis

Cells were fixed with 70% ethanol at -20°C, washed with PBS and incubated for 15 minutes at room temperature in PBS containing 10U ml $^{-1}$  RNaseA and 30 $\mu$  ml $^{-1}$  propidium iodide. Samples were analysed on a FACSCalibur flow cytometer (Beckton Dickinson) using CellQuest software.

Results

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#### Example 1

# Inhibition of SIRT expression by RNAi

A siRNA specific to human SIRT1 was designed which had no more than ~60% homology to any other human sirtuin cDNA (Figure 1A).

BLAST searching of human genomic and EST databases reported only the SIRT1 gene itself as a significant match. As a positive control we used an siRNA directed to lamin A/C (Dharmacon; Elbashir et al., 2001) (Fig 1B); as a negative control we used an siRNA directed to the BCR-ABL fusion oncogene (MWG-Biotech) which is biologically active only in a background of BCR-ABL (Fig 1C). The siRNAs were introduced to wild-type or p53-/- or Bax-1- HCT116 colorectal cancer cells by cationic-based lipid transfection (Oligofectamine TM, Invitrogen) according to the manufacturer's instructions. Cells were harvested 48 hours post-transfection for the preparation of total cellular RNA and at 12 hour intervals post-transfection for the preparation of protein lysates and protein lysates.

One-step RT-PCR showed that only the SIRT1 siRNA reduced the abundance of SIRT1 mRNA (Fig 2B); the mock transfections and control siRNAs did not alter SIRT1 mRNA. RT-PCR specific for the lamin A mRNA demonstrated that the lamin A/C siRNA was active in HCT116 cells (Fig 2D), and therefore that an active siRNA process does not alter SIRT1 mRNA.

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Additionally, silencing of SIRT1 mRNA did not alter lamin A mRNA, and none of the siRNAs altered abundance of the mRNAs chosen as controls, vimentin (Fig 2C) and GAPDH (Fig2A). These data support the specificity of the SIRT1 siRNA for RNAi.

Western blots were performed to establish the efficacy of RNAi in inhibiting protein expression. Figs 3A and 7A/B show that only LaminA/C siRNAs silence Lamin A protein expression in  $p53^{+/+}$  cells (top) and  $p53^{-/-}$  cells under optimal growth conditions, but silencing of SIRT1 can downregulate LaminA if cellular stress is applied. (Note that the antibody used does not seem to recognise

Lamin A in whole cell lysates). Fig 7C/D confirms that BCR-Abl siRNA does not induce RNAi in cells lacking BCR-ABL. Fig 3 demonstrates the efficacy of SIRT1 siRNA in reducing SIRt1 protein expression.

Significant silencing at the protein level was observed for LaminA/C which had not been indicated by the RT-PCR. This suggests that the relationship between silencing at the mRNA and protein levels is not linear. Likewise, blots probed for SIRT1 showed a profound silencing at the protein level (Figure 3) although RT-PCR had shown only ~50% mRNA degradation. Possibly this reflects protection or sequestration of a pool of the mRNA, such that it is not accessible to components of the RNAi machinery.

### Example 2

Effect of SIRT1 siRNA on p53 stability and activity

We went on the investigate the effect of SIRT1 siRNA on p53
stability and activity. In response to cellular stress, it
appears that stabilisation of p53 requires only its prior
destabilisation by HDM2 (Blattner et al., 1999). The most
significant post-translational modification in this regard is
phosphorylation of serine 20, now generally accepted as the
major mechanism through which p53 is removed from HDM2-mediated
negative regulation (Chehab et al., 1999; Dumaz et al., 2001).
Phosphorylation of serine 15, previously thought to be involved
in stabilisation of p53 (Shieh et al., 1997; Unger et al.,
1999), is now believed to be primarily involved in the
activation of p53 as a transcription factor (Dumaz and Meek,
1999). The acetylation of the carboxy-terminus of p53 occurs in
response to most stimuli that stabilise p53 (Itoh et al., 2001).

Although the function of acetylation of p53 is not fully understood it appears to be stimulated by phosphorylation of serine 15 (Dumaz and Meek, 1999; Lambert et al., 1998), implicated in nucleotide excision repair (Rubbi and Milner, 2003) and in the recruitment of acetylase enzymes to chromatin (Espinosa and Emerson, 2001), and required for efficient p53dependent apoptosis (Luo et al., 2001; Vaziri et al., 2001). Although it has been demonstrated that, under certain circumstances, neither amino-terminal phosphorylation nor carboxy-terminal acetylation are obligate events for the stabilisation and activation of p53 (Blattner et al., 1999; Ashcroft et al., 2000), the weight of evidence supports a general model whereby stress stimuli signal to p53 via phosphorylation of amino-terminal serine residues. These phosphorylations stabilise p53 and promote interaction with acetylase enzymes; acetylation of the carboxy-terminus fully activates p53 as a transcription factor and may help stabilise p53 (Nakamura et al., 2000).

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To determine if silencing of SIRT1 could stabilise p53, we performed Western blots using the pantropic anti-p53 antibody DO-1 in cells treated with SIRT1. This revealed that silencing of SIRT1 led to a significant stabilisation of p53 (Fig 4) compared to the mock transfected control. It is possible that the RNAi process itself constitutes a low-level stress signal, as a small stabilisation of p53 was observed for the lamin A/C siRNA but not the BCR-ABL siRNA. Interestingly, the p53 stabilised by the SIRT1 siRNA was partitioned between the soluble (cytoplasmic and nucleoplasmic) fraction and the insoluble fraction protein, thought the p53 stabilised by LaminA/C silencing was found only in the soluble fraction.

We then examined in more detail the stabilisation and activation of p53 in HCT116 cells transfected with SIRT1 siRNA. We chose to examine a four-point time course, with sampling at 12, 24, 36 and 48 hours post-transfection. Protein lysates and fractions were prepared and blotted with antibodies to p53 and phosphoSer15 (Figs 4 and 5).

Significant stabilisation of p53 was not observed in whole cell lysates until 36 hours post-transfection, although the soluble fraction showed that there was stabilisation at 24 hours (Fig 4). Intriguingly, stabilised p53 was not detected in the insoluble fraction until 36 hours post-transfection, and was not highly elevated until 48 hours.

- To analyse the activation of p53 as a transcription factor we examined phosphorylation of serine 15. We determined that stabilised p53 was not subject to serine 15 phosphorylation prior to 36 hours post-transfection. Furthermore, that serine 15 phosphorylation was restricted to p53 in the soluble fraction, as no phosphorylation of this residue was detected at any time point in the insoluble fraction (Fig 5). LaminA/C silencing can induce a very small amount of phosphorylation, and BCR-ABL siRNAs do not stimulate phosphorylation.
- This data is somewhat counter-intuitive, as one might assume that transcriptionally competent p53 would be associated with chromatin. However, these cells were cultured under normal conditions and in the absence of any exogenous stress stimulus. It was therefore possible that although the p53 was stabilised and "activated", additional signals or cofactors not present or accessible to the p53 under these conditions were required to activate gene transcription.

### Example 3

## Effect of SIRT1 siRNA on downstream activity of p53

To investigate this, we analysed expression of the two bestcharacterised p53 responsive genes, p21(CIP1/WAF1) and HDM2.
The results for these were essentially identical, as both showed a strong induction of expression at 36 hours that was absent prior to this time point (Figs 5A and 6). There was therefore a good temporal correlation in this data between stabilisation of p53, "activating" post-translational modification, and the induction of transcriptional targets. This effect was seen only in p53<sup>+/+</sup> cells, indicating that the p21 expression was indeed induced by p53 (Fig 6A). Silencing of SIRT1 does not induce expression of p21 if cellular stress is applied (Fig 7B).

### Example 4

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# SIRT1 siRNA induces apoptosis in tumour cells independently of p53

As p21 was effectively induced, we wondered if these cells were cell cycle arrested. Phase contrast images (Figs 8 and 9) showed a very clear phenotype that was apparent at 36 hours, and pronounced by 48 hours. Close examination of these cells under higher magnification revealed reduction of the cell bodies, extensive membrane blebbing and a granular appearance to the nucleus (Fig 8C). Surprisingly, this effect was seen in p53<sup>-/-</sup> cells (Fig 9) as well as p53<sup>+/+</sup> cells (Fig 8), indicating that the apoptosis observed was not occurring via a p53-dependent pathway, counter to previous findings.

30 As these morphological changes are typical of apoptosis, we analysed control and SIRT1-silenced HCT116 cells by flow cytometry (Figure 11). Propidium iodide staining revealed that

there was no significant change in the cell cycle profile of cells transfected with either the lamin A/C or BCR-ABL siRNAs, as compared to the mock-transfected controls. Cells treated with the SIRT1 siRNA showed a marked decrease of the G1 and G2/M populations and a significant increase in sub-G1 population, consistent with apoptotic chromatin fragmentation.

5

As cells treated with SIRT1 siRNA were clearly undergoing apoptosis, we next considered whether in p53+/+ cells the 10 stabilised p53 induced the expression of components of the proapoptotic machinery. Activation of p53 in HCT116 cells has previously been demonstrated to induce expression of two proapoptotic genes, PUMA and Bax (Yu et al., 2003; Zhang et al., 2000). Neither Bax nor PUMA was induced in SIRT1-silenced 15 cells, as compared to the controls (Fig 12A). Indeed, PUMA appeared to be repressed by transfection of the cells with SIRT1 siRNA. Further, SIRT siRNA induced apoptosis even in p53+/+ cells lacking Bax (Fig 10). These results indicate that SIR1 siRNA can induce apoptosis via a p53-independent mechanism even in  $p53^{+/+}$  cells 20

Two modes of apoptosis are present in human cells: The intrinsic pathway, which functions through mitochondria, and the extrinsic pathway, which functions through the activation of cell death receptors (for review see Green, 1998). PUMA and Bax are both components of the intrinsic pathway, and as neither was induced by the stabilised p53, it was necessary to determine which apoptotic pathway had been activated.

The discovery of cross-talk between the intrinsic and extrinsic pathways (Green, 1998) has been complicated by the discovery that cell type determines the extent of cross-talk. This has led

to the idea that cells fall into one of two types. Type I cells can undergo death receptor induced apoptosis independently of mitochondria; type II cells require mitochondrial involvement for death receptor signalling to induce apoptosis (Scaffidi et al., 1998). Importantly, it has been established that HCT116 cells exhibit the behaviour of type II cells (Deng et al., 2002), which implies that the apoptosis induced by SIRT1 treatment most likely functions through mitochondria.

### 10 Example 5

SIRT1 siRNA does not induce apoptosis in normal cells

We decided to test the effect of SIRT1 siRNA on a non-tumour
cell line. Treatment of primary human normal diploid
fibroblasts (NDFs) with SIRT1 siRNA not only failed to provoke
the massive apoptosis seen in HCT116 cells, it did not appear to
induce apoptosis at all. Cells were still healthy 5 days posttransfection (Fig 13).

We have demonstrated that silencing of SIRT1 in HCT116

20 colorectal cancer cells provokes massive apoptosis. The tumour suppressor p53 is stabilised in response to SIRT1 silencing, and undergoes at least partial activation for transcriptional function. This effect is independent of p53, and we did not detect induction of either PUMA or Bax, two pro-apoptotic genes that have been shown to be critical for p53-dependent apoptosis in colorectal cancer cells. Moreover, apoptosis was not induced by SIRT1 siRNA in normal cells, indicating this effect may be specific to tumour cells.

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#### CLAIMS

- 1. A method of treating a proliferative disease comprising administering to an individual in need thereof an effective amount of a SIRT1 inhibitor.
- 2. A method according to claim 1 wherein the disease is cancer.
- 3. A method according to claim 2 wherein the cancer is a colorectal carcinoma.
- 4. An in vitro method of inducing apoptosis in a cell comprising administering a SIRT1 inhibitor to said cell.
- 5. A method according to claim 4, wherein the cell lacks at least one of functional p53, Bax and PUMA protein.
- 6. A method according to claim 4 or claim 5 wherein the cell is a tumour cell.
- 7. A method according to any one of claims 1 to 6, wherein the SIRT1 inhibitor is a siRNA, a dsRNA, a nucleic acid encoding such RNA, or a SIRT1 antisense RNA.
- 8. A SIRT1 inhibitor for use in a method of medical treatment.
- 9. A SIRT1 inhibitor for use according to claim 8, wherein said treatment is treatment of a proliferative disease.

- 10. A SIRT1 inhibitor for use according to claim 8 or claim 9 which is a siRNA, a dsRNA, a nucleic acid encoding such RNA or a SIRT1 antisense RNA.
- 11. Use of a SIRT1 inhibitor in the manufacture of a medicament for the treatment of a proliferative disease.
- 12. Use according to claim 11, wherein the proliferative disease is cancer.
- 13. Use according to claim 12, wherein the cancer is a colorectal carcinoma.
- 14. Use according to claim 12, wherein the cancer cells lack at least one of functional p53, Bax and protein.
- 15. Use according to any one of claims 11 to 14, wherein the SIRT1 inhibitor is a siRNA, a dsRNA, a nucleic acid encoding such RNA or a SIRT1 antisense RNA.
- 16. A siRNA which inhibits expression of SIRT1 in a cell.
- 17. A siRNA according to claim 16 which comprises a contiguous sequence of 10-30bp from the sequence of SEQ ID No. 2.
- 18. A siRNA according to claim 17 which is between 19 and 22 bp in length.
- 19. A siRNA according to claim 18 which is 19bp in length.

- 20. A siRNA according to claim 19 which has the siRNA sequence of SEQ ID No 1.
- 21. A composition comprising a siRNA according to any one of claims 16 to 20 and a pharmaceutically acceptable excipient.
- 22. The method of any one of claims 1 to 6, wherein the SIRT1 inhibitor is a siRNA according to any one of claims 16 to 20.
- 23. The use of any one of claims 11 to 14, wherein the SIRT1 inhibitor is a siRNA according to any one of claims 16 to 20.
- 24. A method of identifying a SIRT1 inhibitor for use in a method according to any one of claims 1 to 7, the method comprising
- administering a candidate compound to cultured tumour cells in vitro;
- determing whether SIRT expression and/or activity is reduced in said cells; and
- assaying for apoptosis of said cells.
- 25. A method according to claim 24, wherein the cells lack at least one of functional p53, Bax and protein.
- 26. A method according to claim 24 or claim 25, further comprising the steps of administering said candidate compound to cultured normal cells in vitro and assaying for apoptosis of said cells.



Figure 1

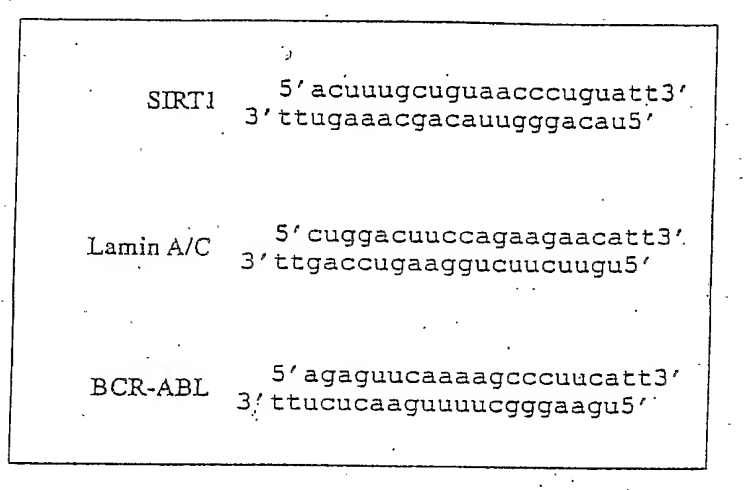
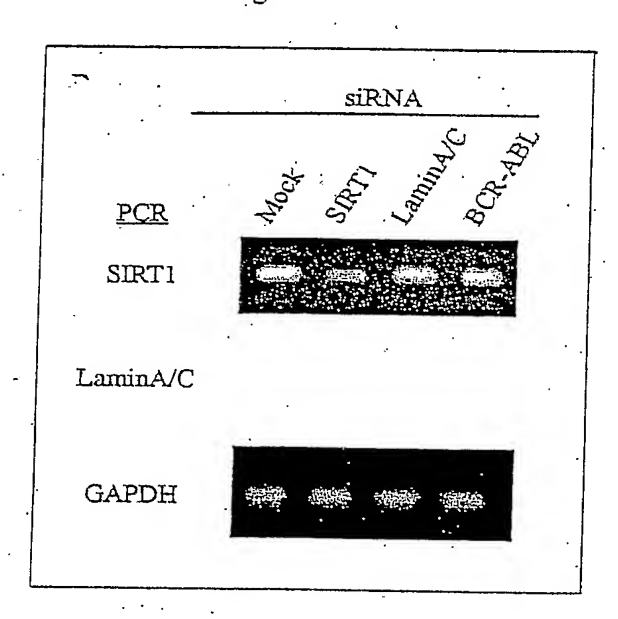


Figure 2



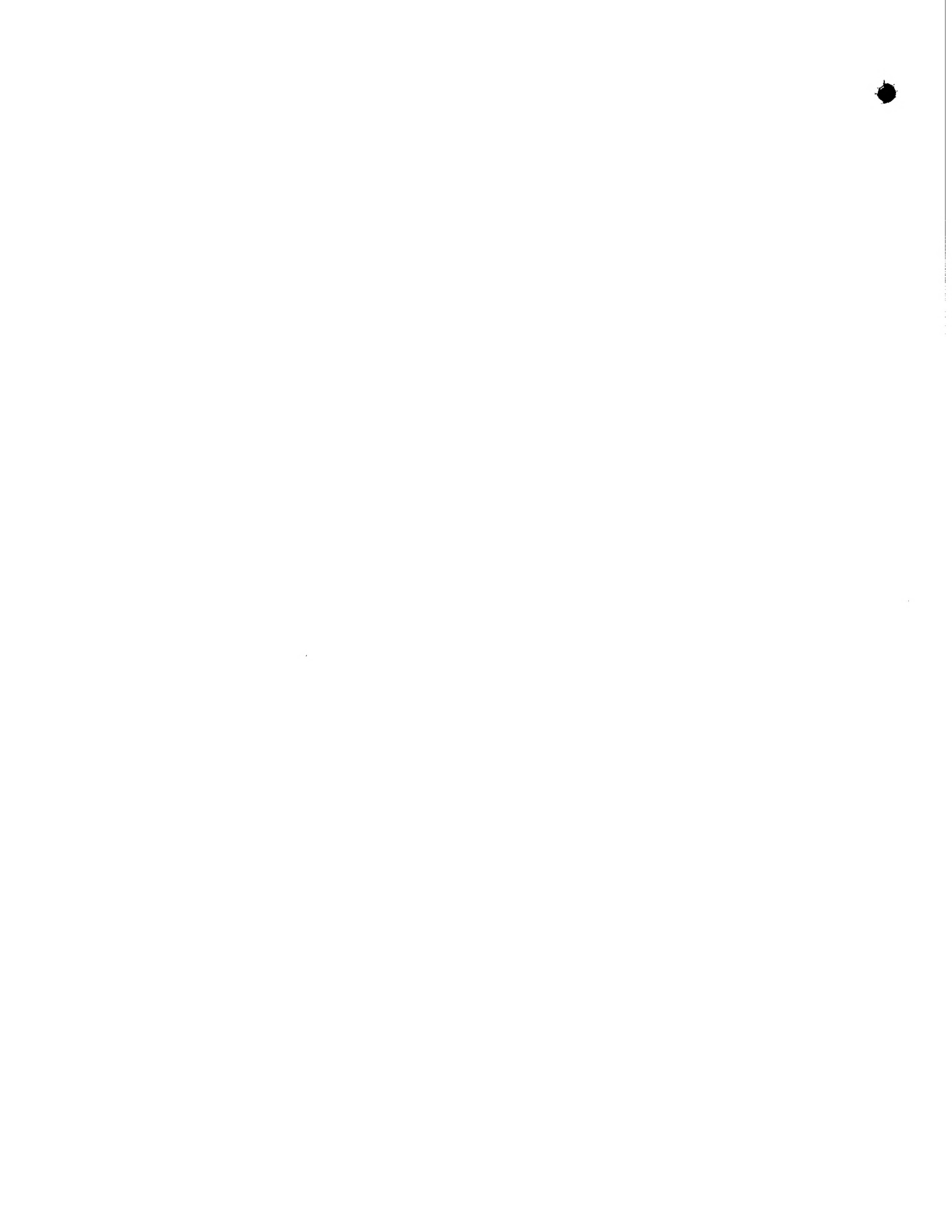
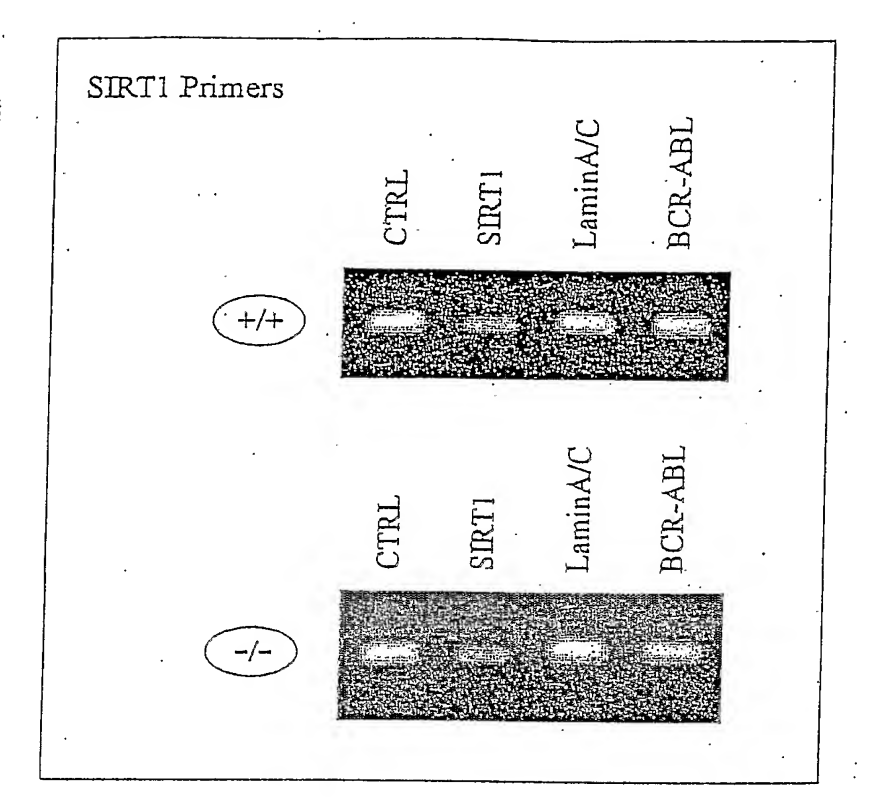
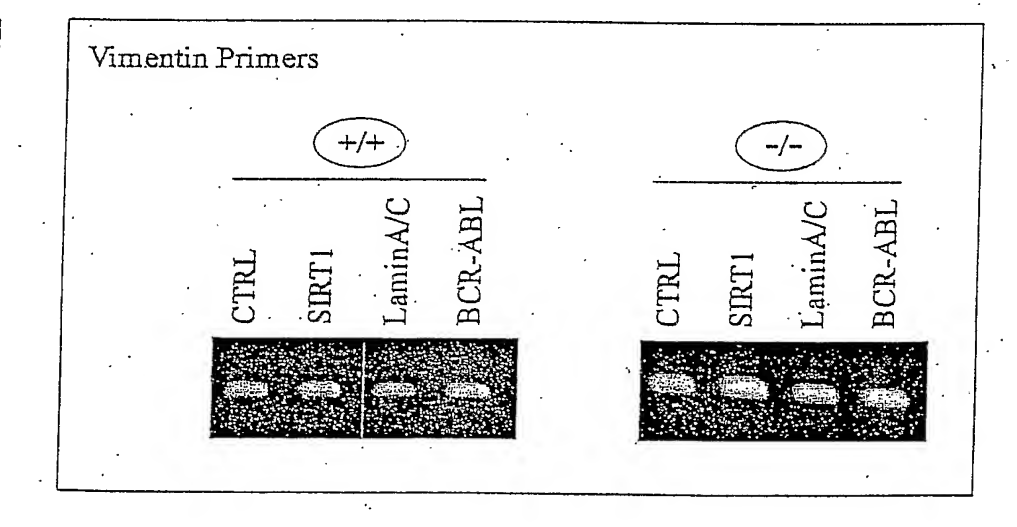


Figure 2

B



C



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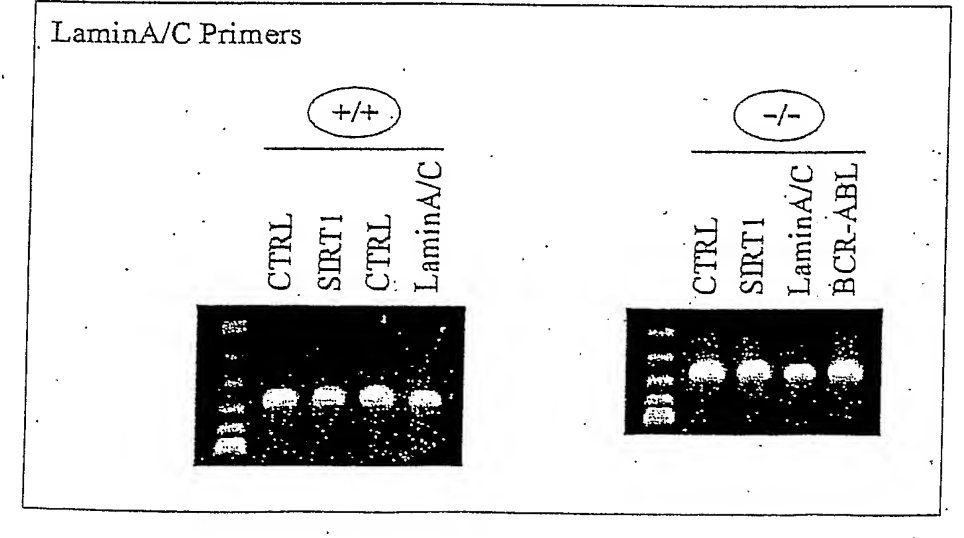
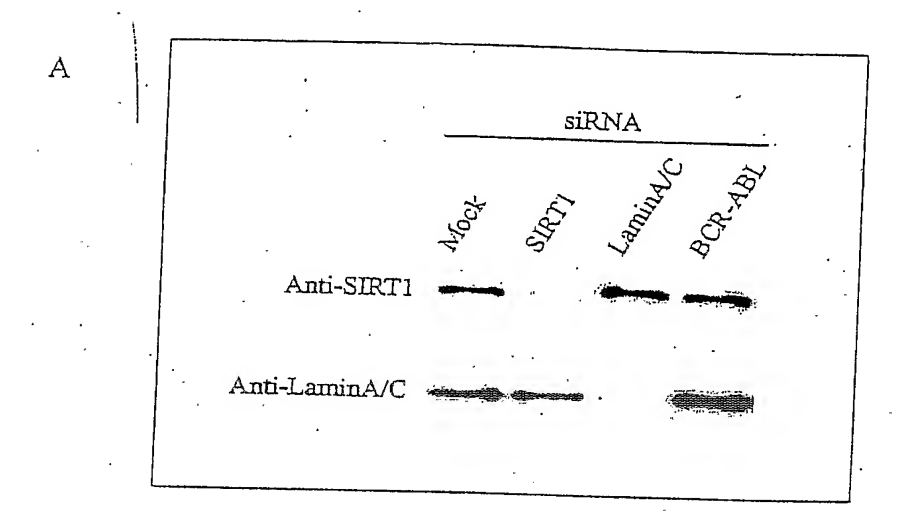
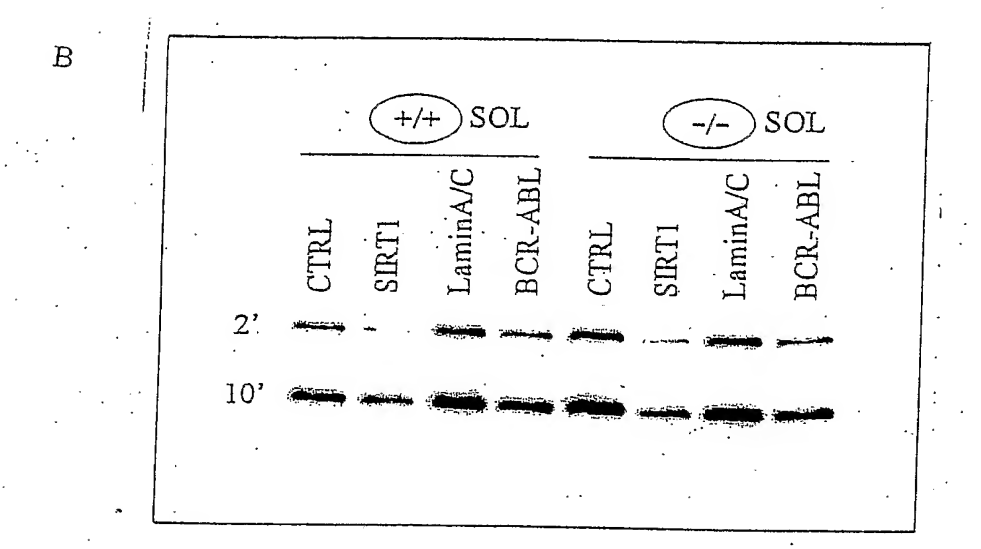
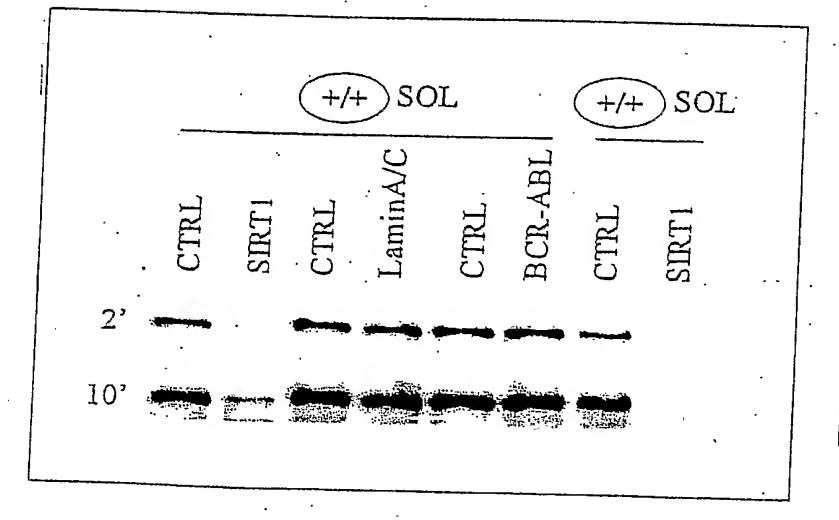
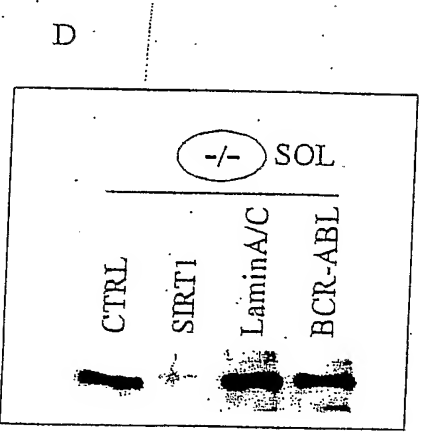


Figure 3



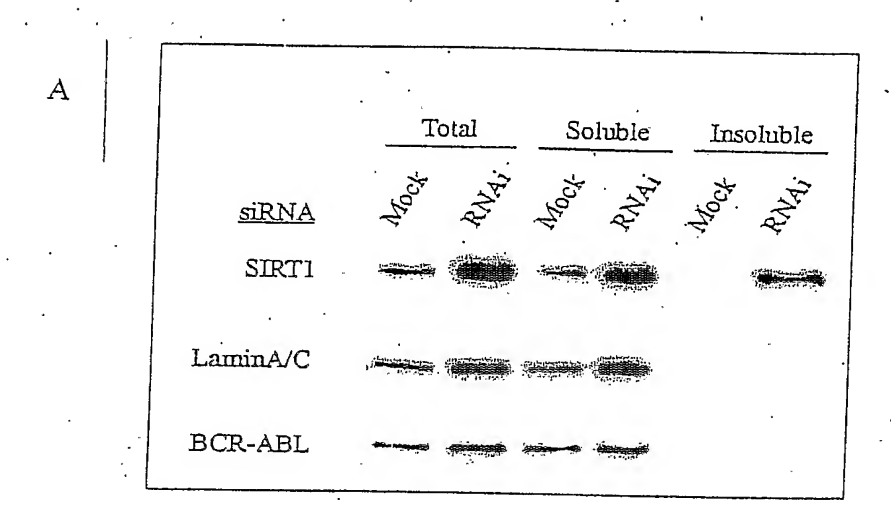


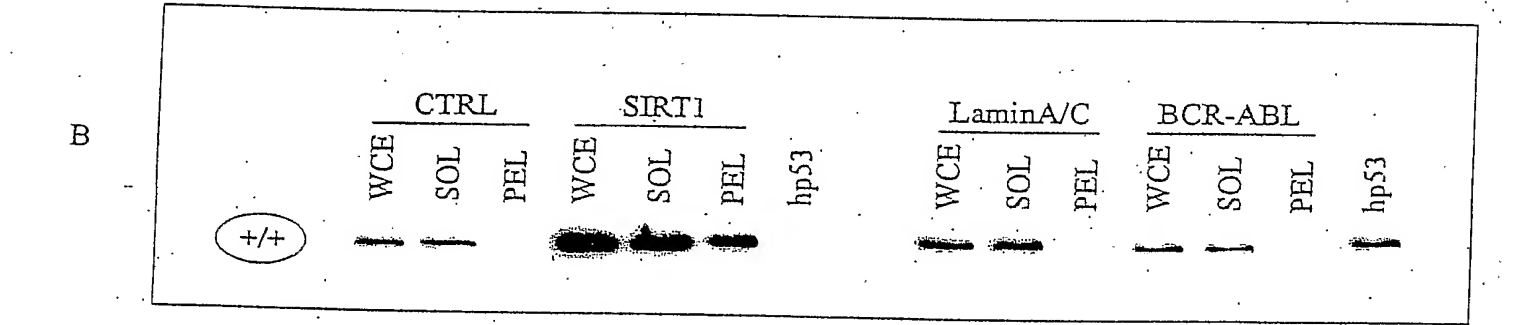


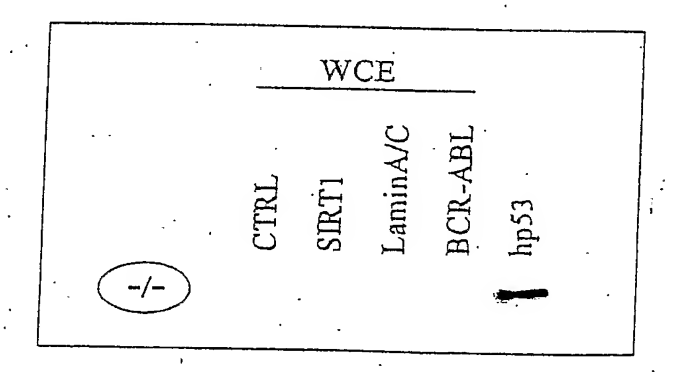


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Figure 4







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Figure 4

•	CTRL	NCE ITAIS	CTRL	SIRT1		SIRT1	hp53
		Mina					
٠		* ~~~		•	•		
٠	•	A/C = 3.7		A/C TO	<u>P</u> .	A/C TE	•
	CTRL	LaminA/C	CTRL	LaminA/C	CTRL	LaminA/Ç	hp53
•	وره معامله و مطوعه میوند. به متا میشد و چی هستانند. به متا میشد و چی هستانند.						
	W(	CE	SC	OT.	PE	77	
•	CTRL	BCR-ABL	CTRL	BCR-ABL	CTRL	BCR-AB	hp53

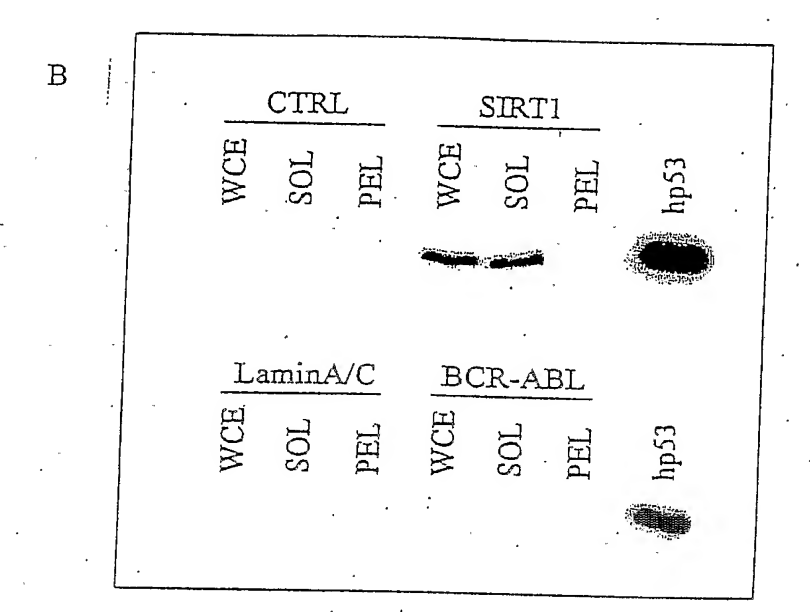


Figure 5

	***************************************	·	<b>lock</b>			SIRT	l siRNA		
Protein	12	24	36	48	12	. 24		48	Antibody
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Insoluble				پېښو د بختر در منځس					
		······································	ock	:		· SIRT	siRNA	·	
	12	24	36	48	12	· 24	36	48	•
Total	-		•		-				•
Soluble	•			•	•		ور المتلافين والمتلافق	to the same of	Anti-phosphoserine
T7 1 7	•	•	•	·					rand-phosphoseime
Insoluble								••	•   .
		Mo	; ck	·	7	SIRTI	siRNA		1.
	12	24	36	48	12	24	36	48	-
Total			¢ 3000			· <del>····</del> -		70	Anti-p21
Total									



Figure 5



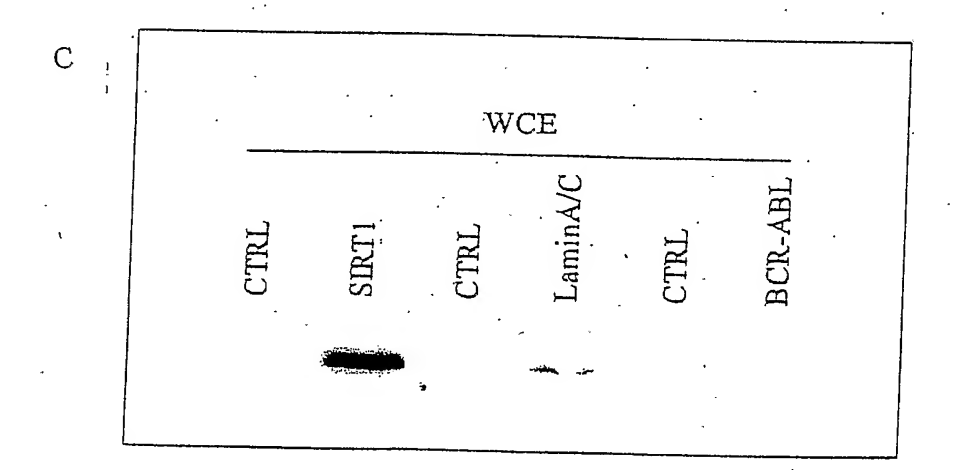
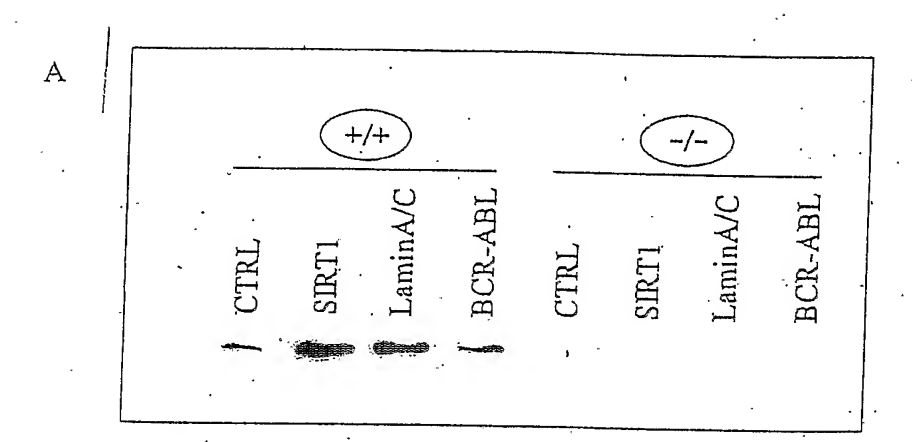


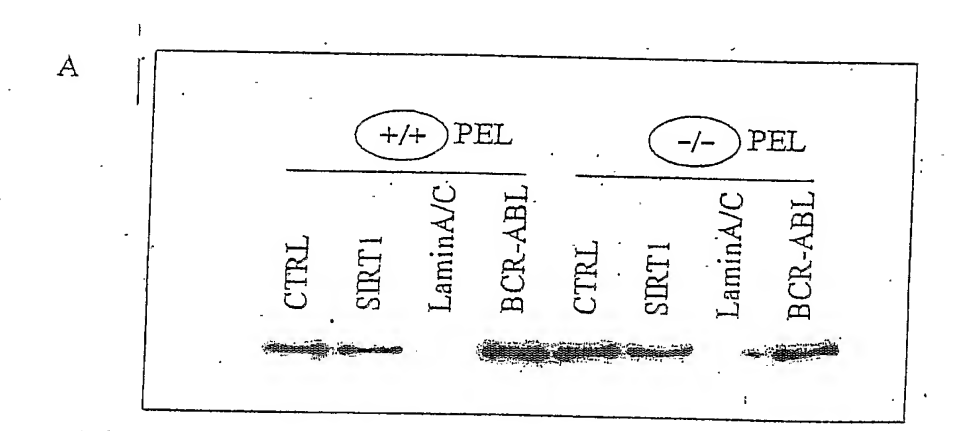
Figure 6

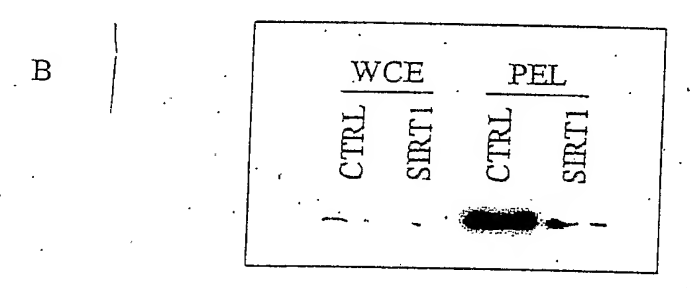


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Figure 7





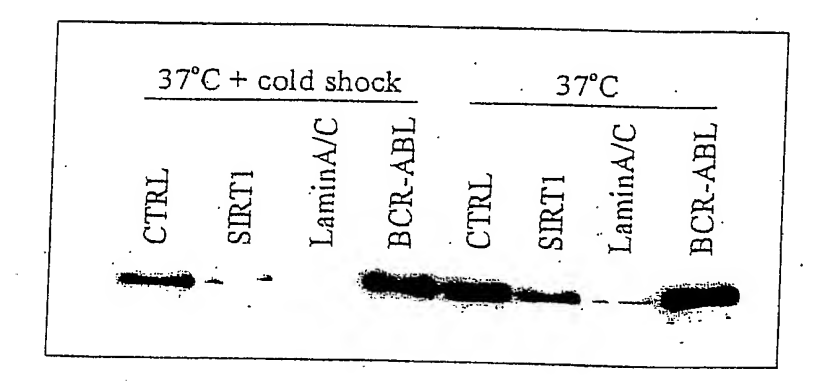


Figure 7

STRT1

LaminA/C

BCR-ABL

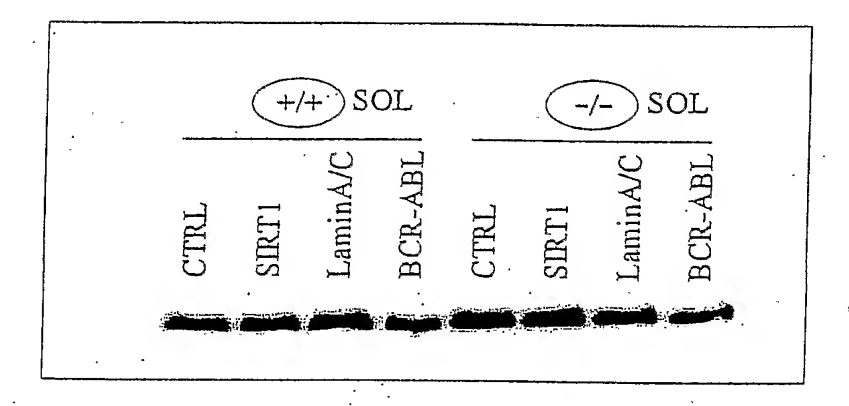
STRT1

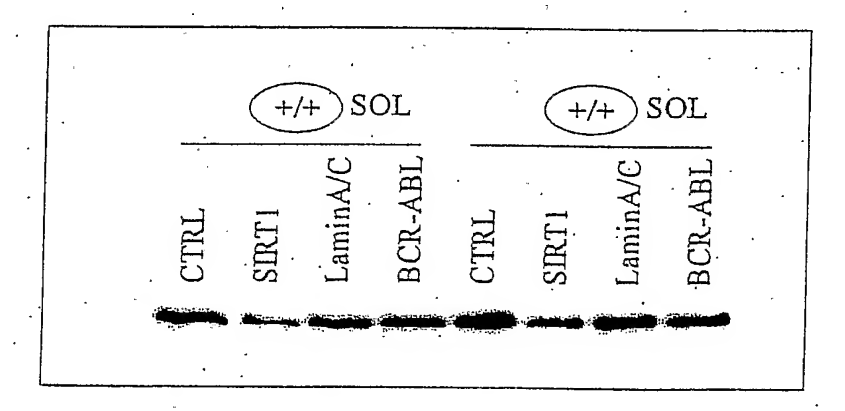
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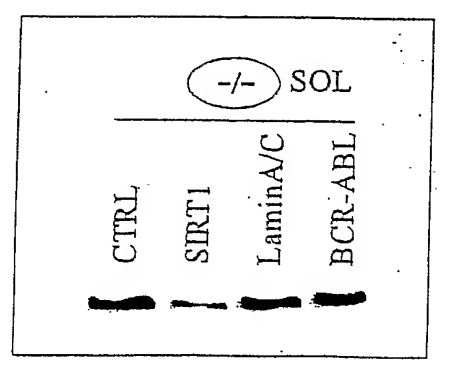
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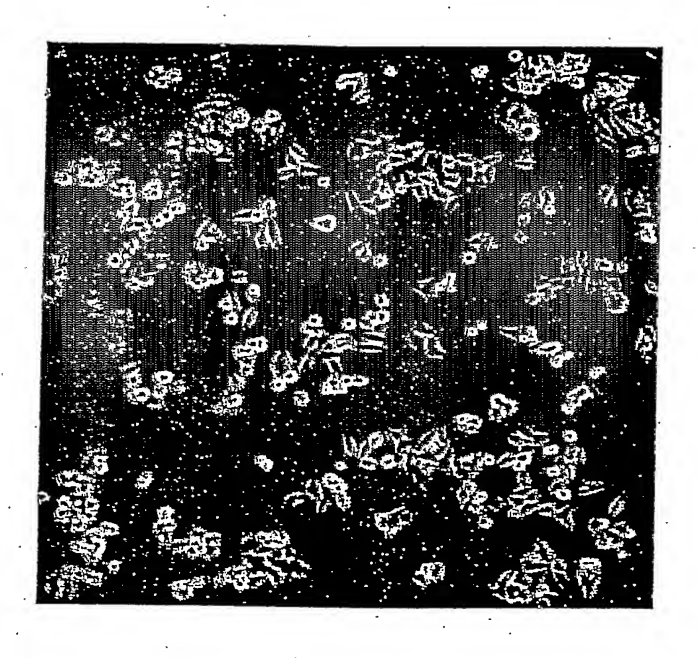
STRT1

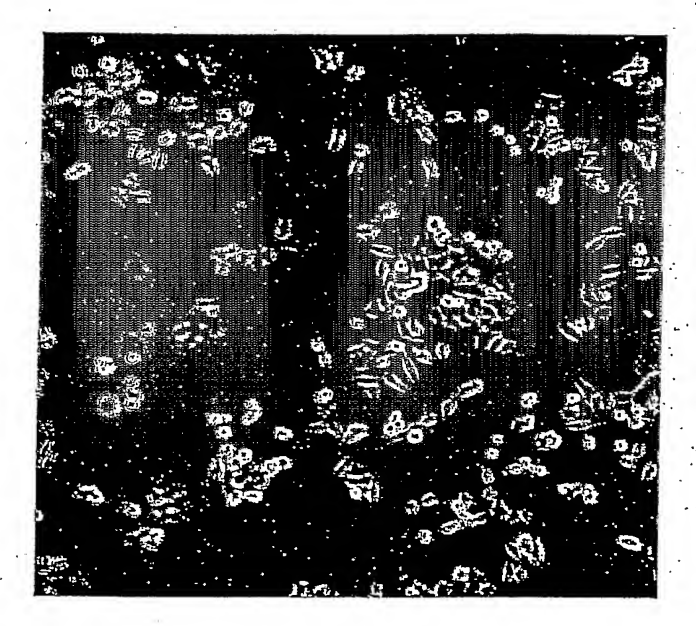


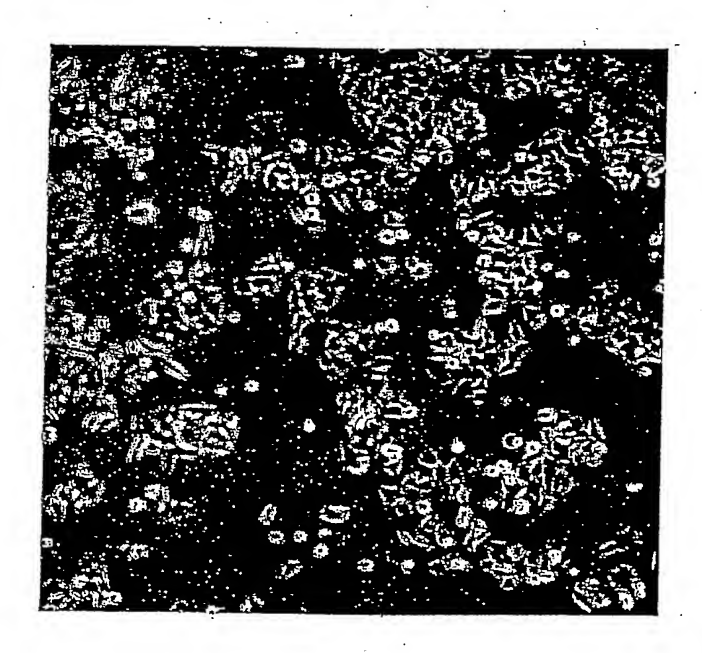


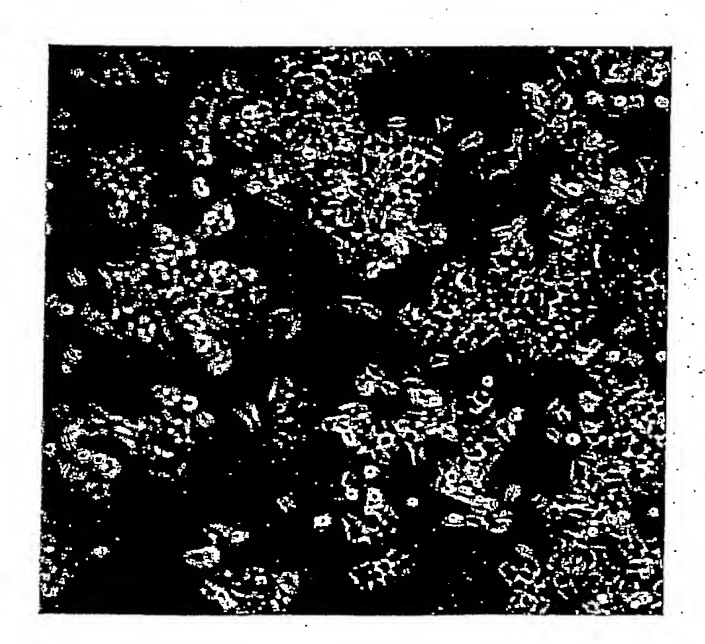




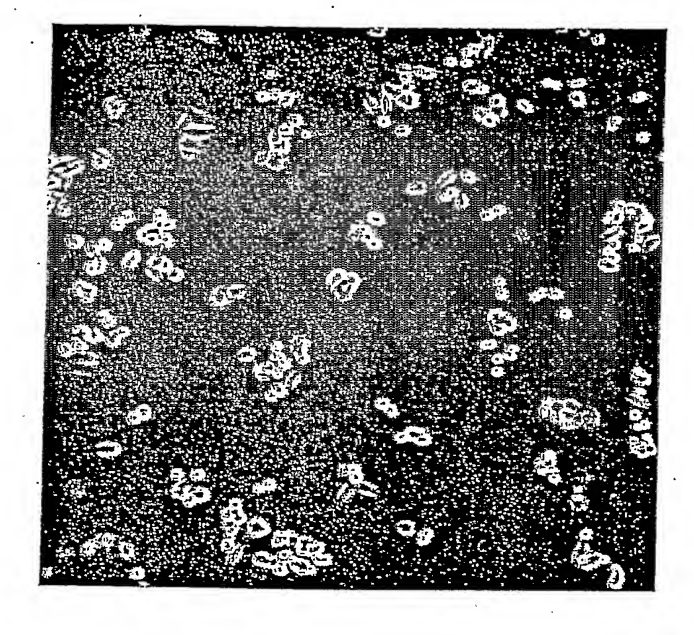


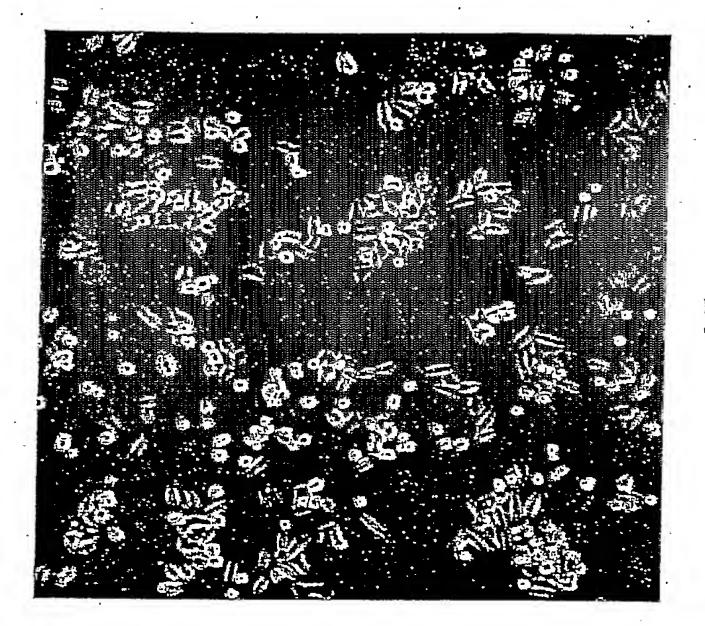


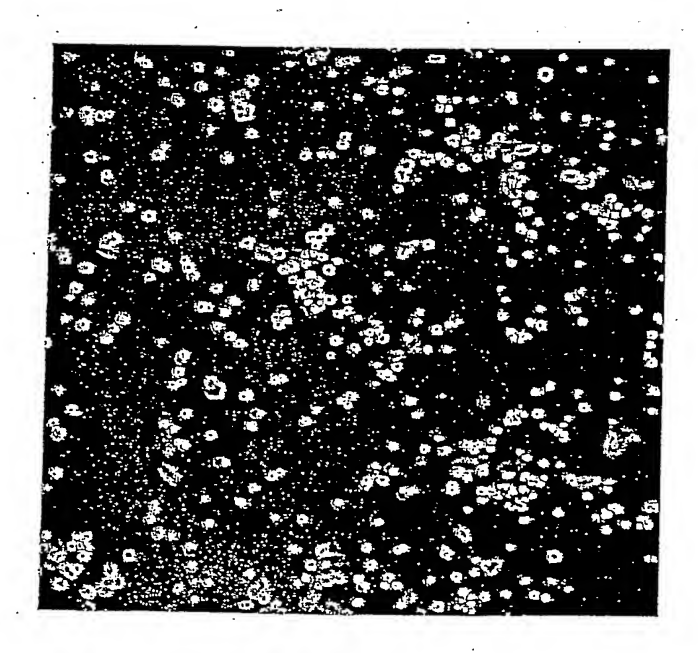


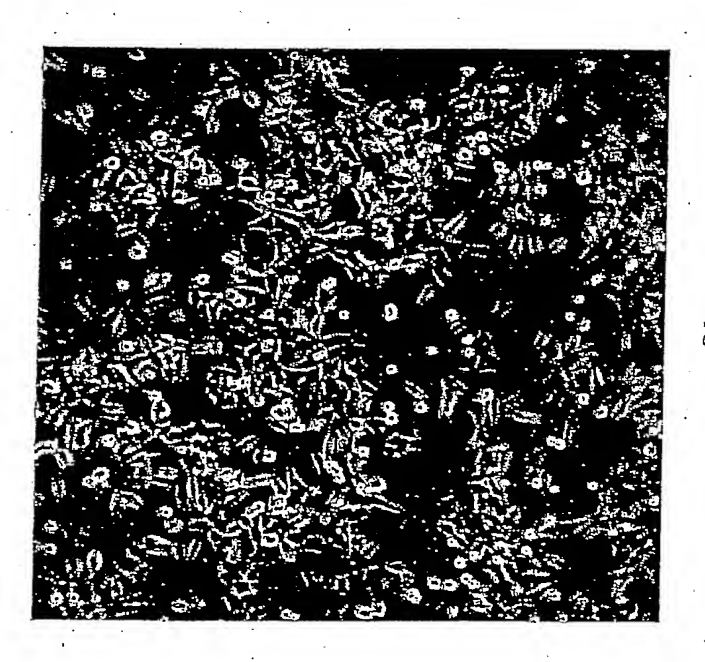






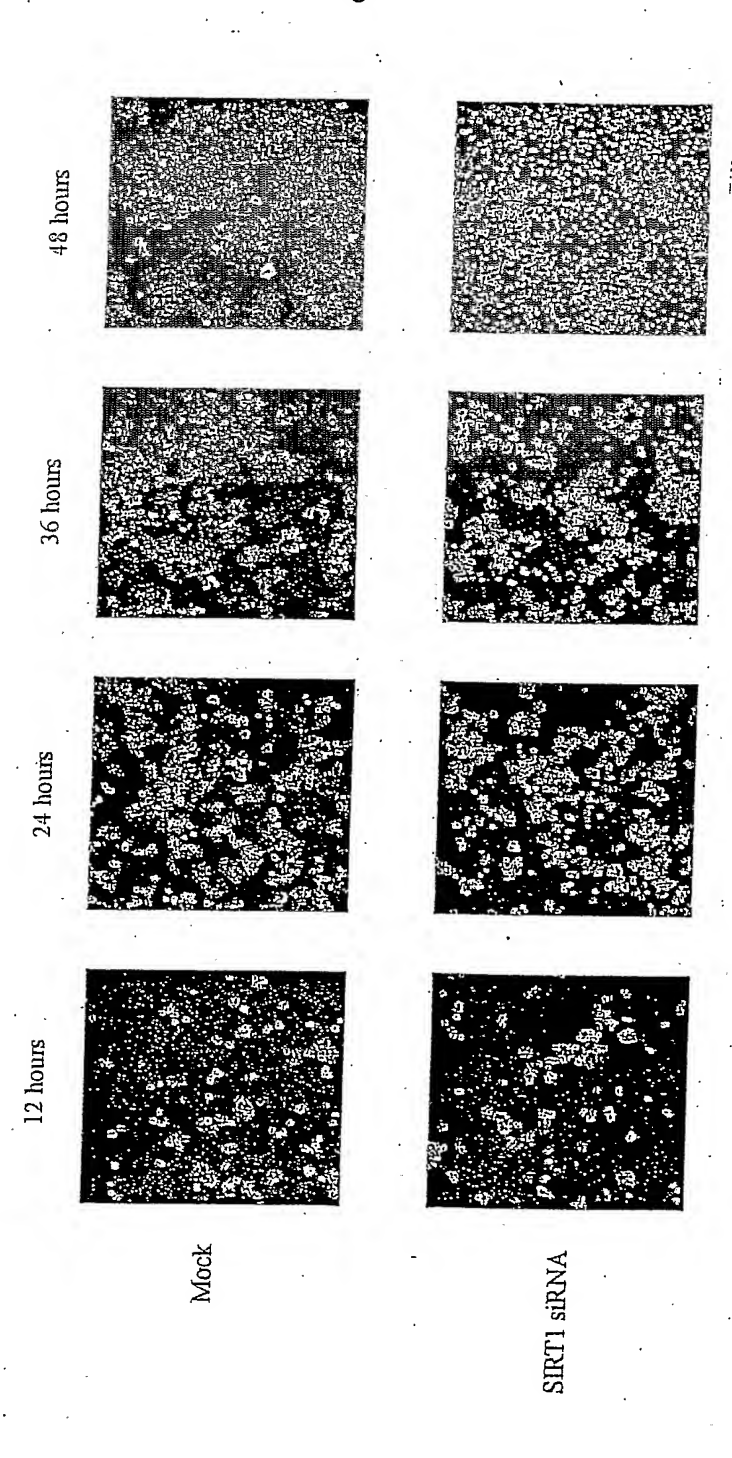


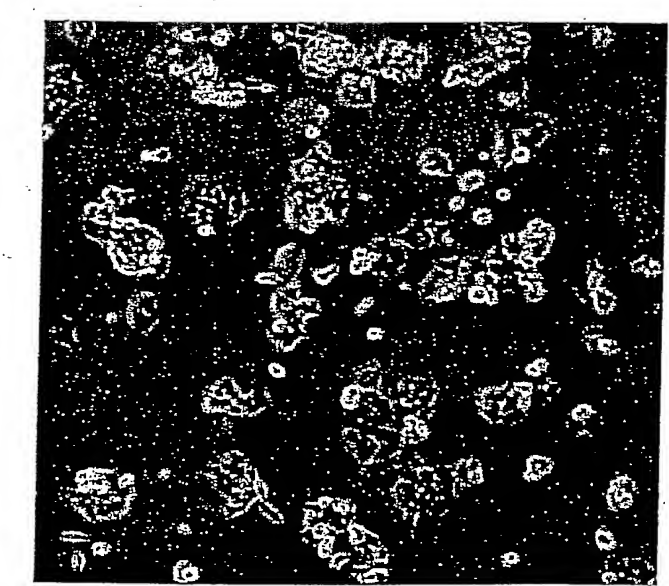


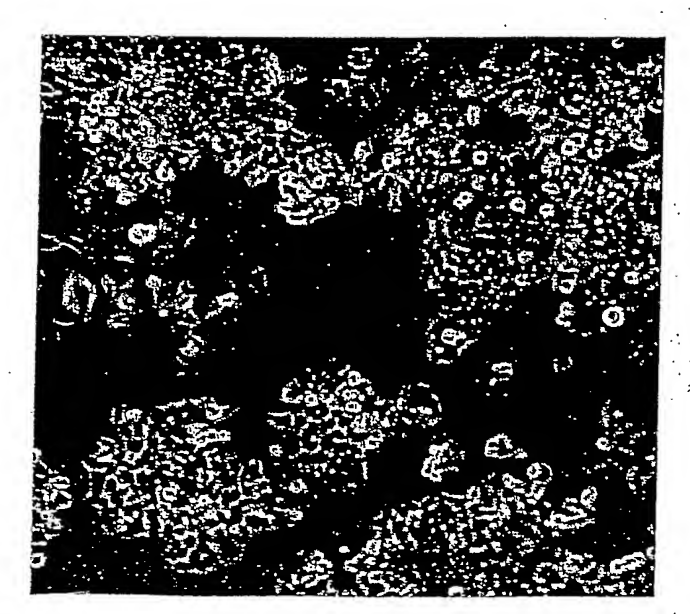


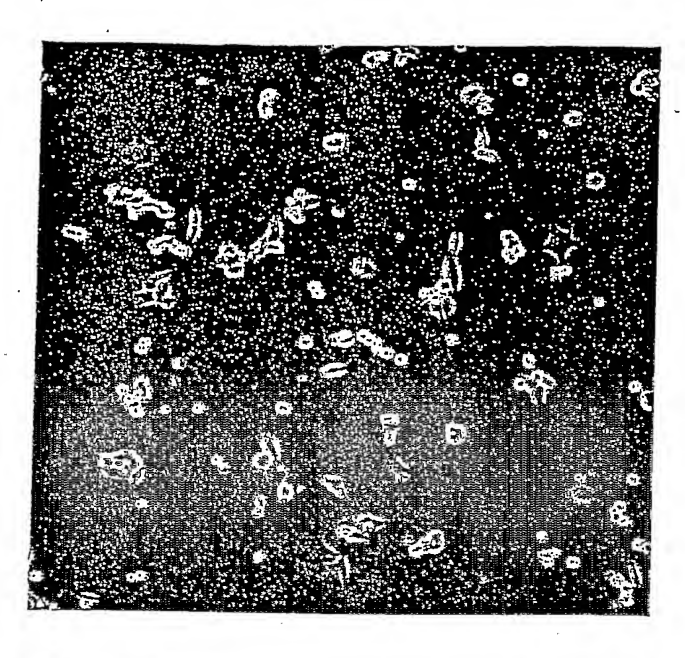
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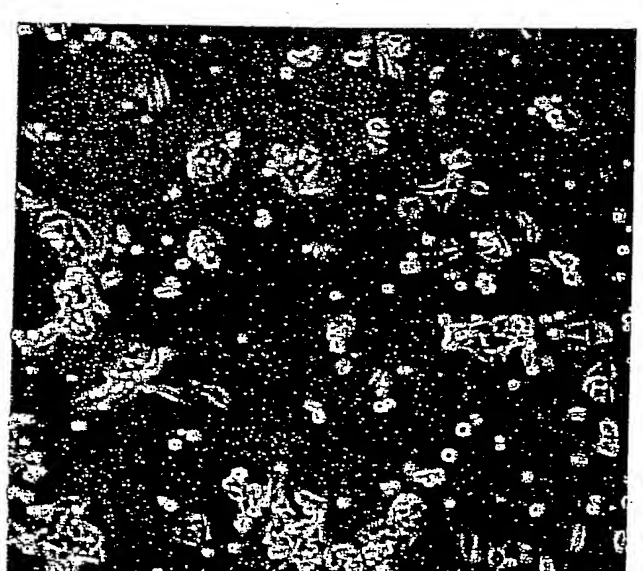
Figure 8B

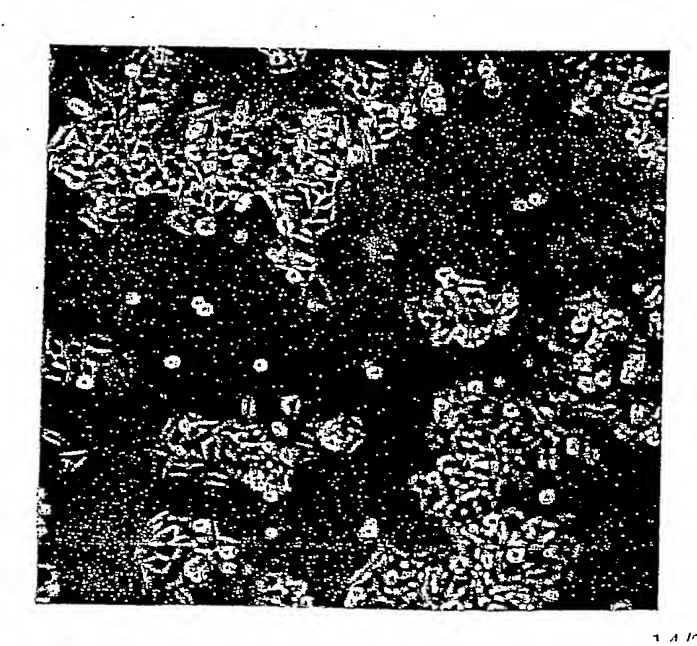












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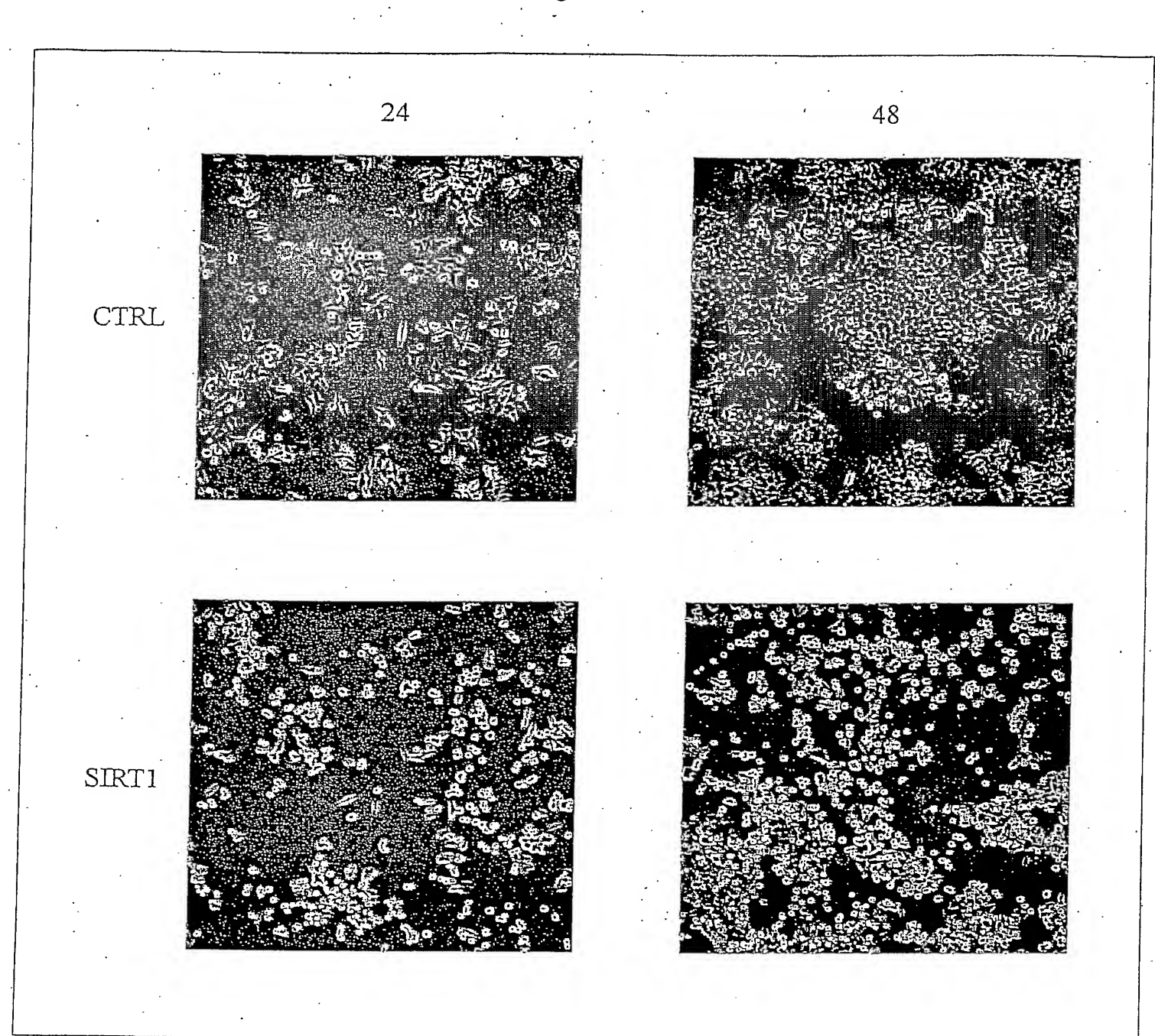
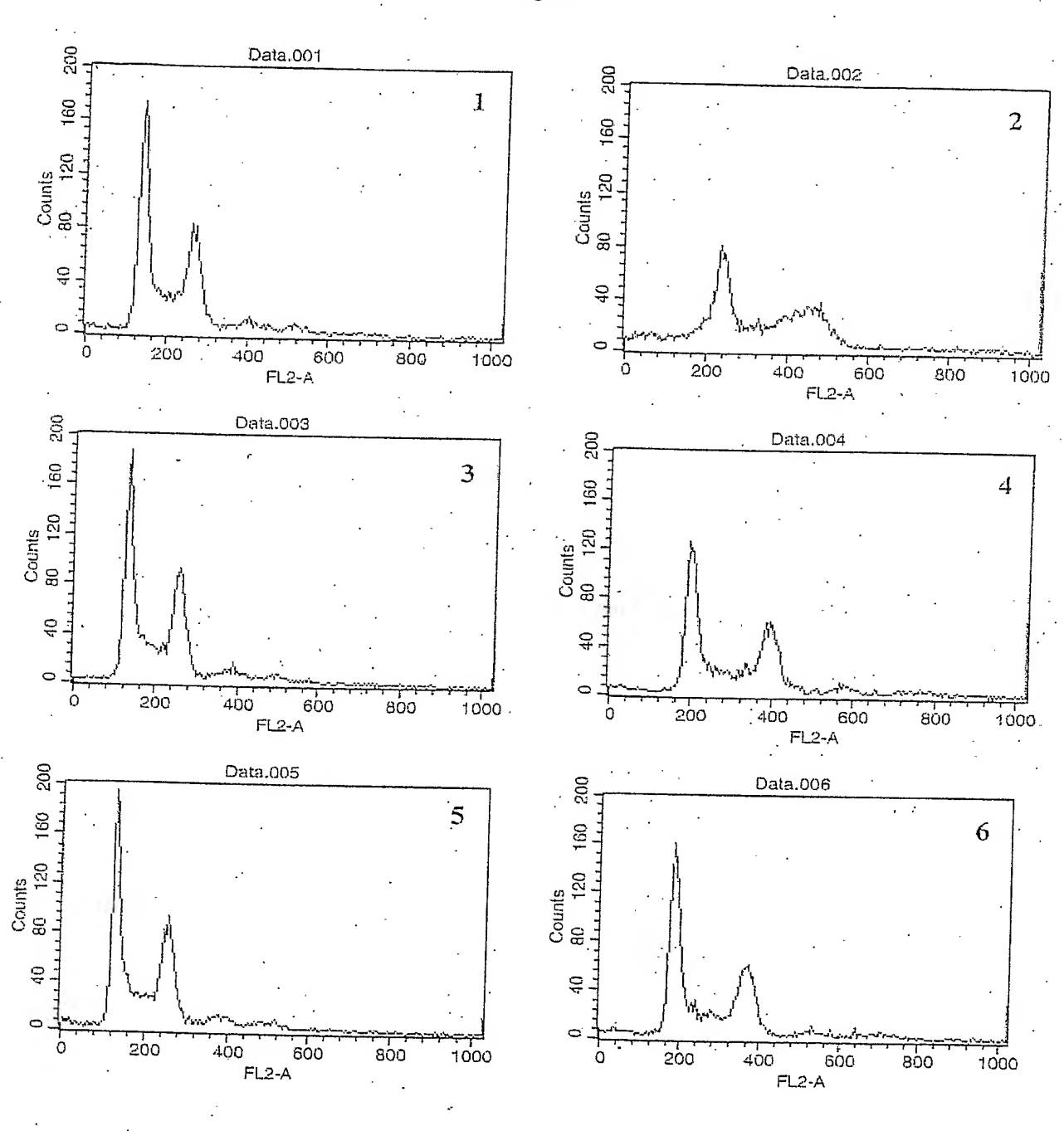




Figure 11



- 1 Mock
- 2 SIRTI siRNA
- ·3 Mock
- 4 LaminA/C siRNA
- 5 Mock
- 6 BCR-ABL siRNA



Figure 12

•			Mock	,		SIRT	l siRNA	•
	12	24	36	48 ·	. 12 ·	24	36	48
Anti-Bax	,	- Minne				·	-	الاسب
Anti-PUMAα		Marie Control					.metata di Maria	· ·



Figure 13

## NDF: Control [Mock] Transfection

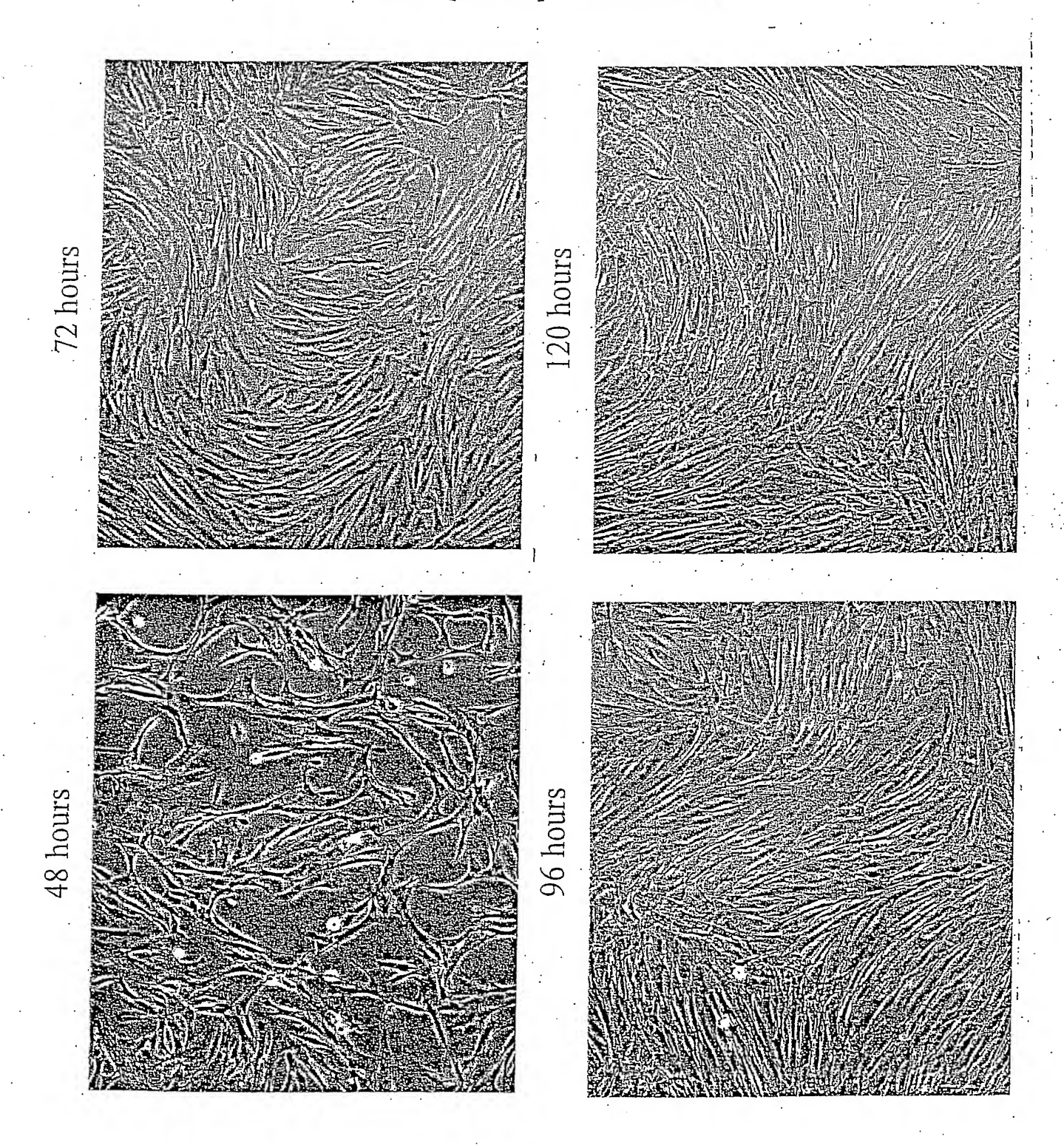
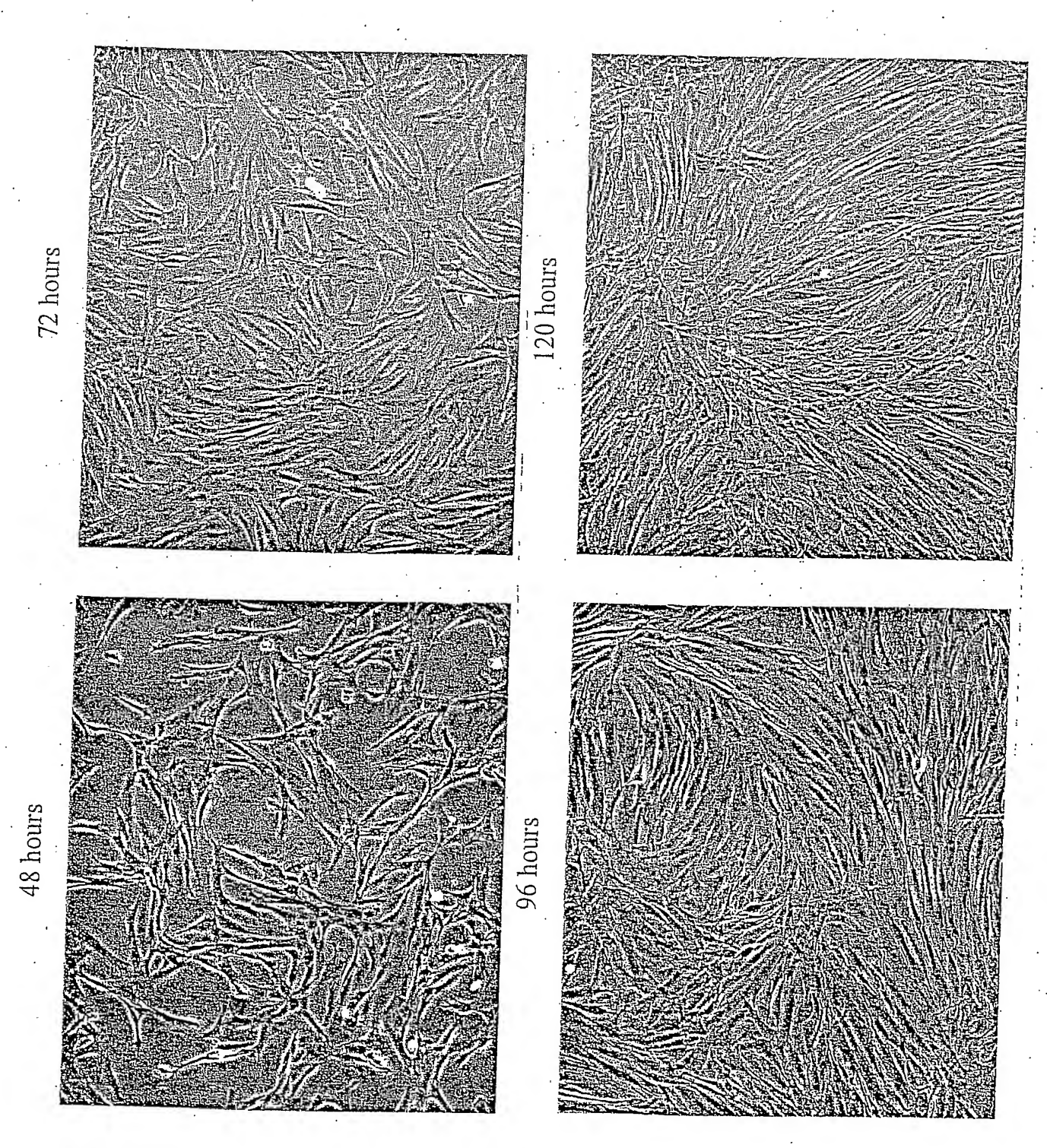


Figure 13

# NDF: SIRT1 siRNA Transfection



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## Figure 14 (1)

	•	-				•
		1 ATGGCGGA	ACG AGGCGGCC	CT CGCCCTTC	AG CCCGGCGG	CT CCCCCTCGG
		TACCGCCT	GC TCCGCCGG	GA GCGGGAAG	TC GGGCCGCC	GA GGGGGAGCC
				•		
	5	1 GGCGGGGG	CC GACAGGGA	GG CCGCGTCG	TC CCCCGCCG	GG GAGCCGCTCC
		CCGCCCCC	GG CTGTCCCT	CC GGCGCAGC	AG GGGGCGGC	CC CTCGGCGAGG
		•	•		•	•
	101	L GCAAGAGG(	CC GCGGAGAG	AT GGTCCCGG	CC TCGAGCGGA	G CCCGGGCGAG
		CGTTCTCC	GG CGCCTCTC	TA CCAGGGCC	GG AGCTCGCCI	C GGGCCGCTC
		•	•	ФY	а	•
	151	- 444667	GG CGGCCCA	GA GCGTGAGG:	rg ccggcggcg	G CCAGGGGCTG
		GGGCCACCC	CC GCCGGGGT	CT CGCACTCC	AC GGCCGCCGC	C GGTCCCCGAC
	201					•
	201	0000000000	CG GCGGCGCC	GG CGCTGTGG(	CG GGAGGCGGA	.G. GCAGAGGCGG
	Ŷ	GGGCCCACG	gc cgácácca	CC GCGACACC	SC CCTCCGCCT	C CGTCTCCGCC
	251	· CCCCCCCAC				
	201	999996119	GGGGGGGGGG	AA GAGGCCCAG	G CGACTGCGG	C GGCTGGGGAA
		GCCGCCGIC	of Gelectical	M CTCCGGGTC	C GCTGACGCC	G CCGACCCCTT
	301	GGAGACAAT	re eccecece			
		CCTCTGTTA	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	T GCAGGGCCC	A TCTCGGGAG	C CACCGCTGGC
	•	·	ic coggeogg	. CGICCCGGG	T AGAGCCCTC	G GTGGCGACCG
	351	CGACAACTT	G TACCACCAA	G ACGACGACC	· · · · · · · · · · · · · · · · · · · ·	G GAGGAGGAAG
		GCTGTTGAA	C ATGCTGCTT	O TGCTGCTGC	T CCTCCCCCT	CICCICCTIC
				0 100100100	T GCTCCCGCTC	CICCICCTTC
	401	· AGGCGGĊGG	C GGCGGCGAT	T GGGTACCGA	$G^{'}\Delta \Psi \Delta \Delta C C \Psi \Psi C \Psi$	GTTCGGTGAT
		TCCGCCGCC	G CCGCCGCTA	A CCCATGGCT	C TATTGGAAGI	CAAGCCACTA
						1 CANGCCACIA
•	451	GAAATTATCA	A CTAATGGTT	T TCATTCCTG	T GAAAGTGATG	S AGGAGGATAG
		CTTTAATAGT	T GATTACCAA	A AGTAAGGAC		TCCTCCTATC.
	•	•	•	· ·		•
	.501	AGCCTCACAI	T GCAAGCTCT	A GTGACTGGA	C TCCAAGGCCA	CGGATAGGTC
	*	TCGGAGTGTA	A CGTTCGAGA	r cactgacct	G AGGTTCCGGT	' GCCTATCCAG
	L. L4					·
	551	CATATACTTT	T TGTTCAGCAZ		A TTGGCACAGA	
		GTATATGAAA	A ACAAGTCGTT	r GTAGAATAC:	r AACCGTGTCT	AGGAGCTTGT
	601	7 mm C mm 7 m m 7			:	•
•	90T	ATTCTTAAAG	G ATTTATTGCC	C GGAAACAATA	A CCTCCACCTG	
		TAAGAATITC	C TAAATAACGO	G CCTTTGTTAT	GGAGGTGGAC	TCAACCTACT
	651	T			•	•
	031		G TGGCAGATTO			CCACCAAAAA
		MIACIGIGAC	C ACCGTCTAAC	: AATAATTATA	GGAAAGTCTT	GGTGGTTTTT
	701	· CCAAAAAAAA	G AAAAGATATI			
	, , , , , ,		TTTTCTATAA			GAAATTACTG
			· TTTTCTHTHH	LIAIGTTAAC	TTCTACGACA	CTTTAATGAC
	751	CAAGAGTGCA	AAAAAATTAT	<u>አ</u> ርጥጥራጥ አ አ ረመ		
			•		GGAGCTGGGG	TGTCTGTTTC
				TOUGHTIGH	CCTCGACCCC	ACAGACAAAG
	801	ATGTGGAATA	CCTGACTTCA	CCTCNNCCCN	TGGTATTTAT	COMOCOOMMO '
		TACACCTTAT	GGACTGAACT	ADDDAAD I DD	ACCATAAATA	GCTCGCCTTG
				· COMBLICCCI	ACCALAAATA	CGAGCGGAAC



8	51 CTGTAGACTT CCCAGATCTT CCAGATCCTC AAGCGATGTT TGATATTGAA GACATCTGAA GGGTCTAGAA GGTCTAGGAG TTCGCTACAA ACTATAACTT	
9( 	O1 TATTTCAGAA AAGATCCAAG ACCATTCTTC AAGTTTGCAA AGGAAATATA ATAAAGTCTT TTCTAGGTTC TGGTAAGAAG TTCAAACGTT TCCTTTATAT	
95	TCCTGGACAA TTCCAGCCAT CTCTCTGTCA CAAATTCATA GCCTTGTCAG AGGACCTGTT AAGGTCGGTA GAGAGACAGT GTTTAAGTAT CGGAACAGTC	
100	1 ATAAGGAAGG AAAACTACTT CGCAACTATA CCCAGAACAT AGACACGCTG TATTCCTTCC TTTTGATGAA GCGTTGATAT GGGTCTTGTA TCTGTGCGAC	
105	2T1PHO  T CAGTGTCATG GTTCCTTTGC  1 GAACAGGTTG CGGGAATCCA AAGGATAATT CAGTGTCATG*GTTCCTTTCC	
	CTTGTCCAAC GCCCTTAGGT TTCCTATTAA GTCACAGTAC*CAAGGAAACG	•
. 110:	AACAGCATCT TGCCTGATTT GTAAATACAA AGTTGACTGT GAAGCTGTAC TTGTCGTAGA ACGGACTAAA CATTTATGTT TCAACTGACA CTTCGACATG	•
1151	1 GAGGAGATAT TTTTAATCAG GTAGTTCCTC GATGTCCTAG GTGCCCAGCT CTCCTCTATA AAAATTAGTC CATCAAGGAG CTACAGGATC CACGGGTCGA	
1201		
1251	TTTACCAGAA CAGTTTCATA GAGCCATGAA GTATGACAAA GATGAAGTTG AAATGGTCTT GTCAAAGTAT CTCGGTACTT CATACTGTTT CTACTTCAAC	
1301	ACCTCCTCAT TGTTATTGGG TCTTCCCTCA AAGTAAGACC AGTAGCACTA TGGAGGAGTA ACAATAACCC AGAAGGGAGT TTCATTCTGG TCATCGTGAT	
1351	ATTCCAAGTT CCATACCCCA TGAAGTGCCT CAGATATTAA TTAATAGAGA TAAGGTTCAA GGTATGGGGT ACTTCACGGA GTCTATAATT AATTATCTCT	
1401	ACCTTTGCCT CATCTGCATT TTGATGTAGA GCTTCTTGGA GACTGTGATG TGGAAACGGA GTAGACGTAA AACTACATCT CGAAGAACCT CTGACACTAC	
	~~~~` "\	
1451	TCATAATTAA TGAATTGTGT CATAGGTTAG GTGGTGAATA TGCCAAACTT AGTATTAATT ACTTAACACA GTATCCAATC CACCACTTAT ACGGTTTGAA dTdTUGAA	•
1501	UGCUGUAACC CUGUAdTdT TGCTGTAACC CTGTAAAGCT TTCAGAAATT ACTGAAAAAC CTCCACGAAC ACGACATTGG GACATTTCGA AAGTCTTTAA TGACTTTTTG GAGGTGCTTG ACGACAUUGG GACAU	
	rJF4	

1551	ACAAAAAGAA TGTTTTTCTT				CCTCTTCATG GGAGAAGTAC
1601	TTTCAGAAGA AAAGTCTTCT		CCAGAAAGAA GGTCTTTCTT		AGATTCTTCA TCTAAGAAGT
1651	GTGATTGTCA CACTAACAGT		CCAAGCAGCT GGTTCGTCGA	_ `	ATGATTTAGA TACTAAATCT
1701	TGTGTCTGAA ACACAGACTT		GTATGGAAGA CATACCTTCT		
1751	CTTCTAGGAA GAAGATCCTT		ATTGCTGAAC TAACGACTTG	AGATGGAAAA TCTACCTTTT	
1801			TGGGGAGAAA ACCCCTCTTT		CTTCAGTGGC GAAGTCACCG
1851			GGCCTAATAG CCGGATTATC		CTCGTCTAAT
•		· :	- L C	2T1R	
1901	•		CAGTATCTGT GTCATAGACA		
1951	ATTTTCCATG		ATATTCAGAC TATAAGTCTG		
1951 2001	ATTTTCCATG TAAAAGGTAC CTCTAGTTCT	CGCGACTCCA .	TATAAGTCTG ACAGTGATAG	AGACTTCTAC	TGCAGAATAG CAGAGTCCAA
	ATTTTCCATG TAAAAGGTAC CTCTAGTTCT GAGATCAAGA GTTTAGAAGA	CGCGACTCCA TGTGGCAGTA ACACCGTCAT ACCCATGGAG	TATAAGTCTG ACAGTGATAG	AGACTTCTAC TGGGACATGC ACCCTGTACG AAATTGAAGA	TGCAGAATAG CAGAGTCCAA GTCTCAGGTT ATTCTACAAT
2001	ATTTTCCATG TAAAAGGTAC CTCTAGTTCT GAGATCAAGA GTTTAGAAGA CAAATCTTCT GGCTTAGAAG	CGCGACTCCA TGTGGCAGTA ACACCGTCAT ACCCATGGAG TGGGTACCTC ATGAGCCTGA	TATAAGTCTG ACAGTGATAG TGTCACTATC GATGAAAGTG	AGACTTCTAC TGGGACATGC ACCCTGTACG AAATTGAAGA TTTAACTTCT AGAGCTGGAG	TGCAGAATAG  CAGAGTCCAA GTCTCAGGTT  ATTCTACAAT TAAGATGTTA  GAGCTGGATT
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#### Figure 15

· hSIRT2			
hSIRT3 hSIRT1	•	**************************************	
HOTKIT		MADEAALALQPGGSPSAAGADREAASSPAGEPLRKRPRRDGPGLERSPGEPGGAAPEREV	60
hSIRT2			
hSIRT3	•	-MAFWGWRAAAALRLWGRVVERVEAGGGVGPFQACGCR	37
hSIRT1		$\cdot$ DA A A D C C D C A A A A A A W D D $\cdot$ A D A D $\cdot$ A A A A C C D $\circ$	12
hSIRT2	• .	: MAEP-DPSHPLETQAG-	15
hSIRT3	•	LVLGGRDDVSAGLRGSHGARGEPLDBARPLORPPRP	73
hSIRT1	•	Y DE DIDDE CEEEE EE VAAAA TOVD DAT EE DODE EE EE EE EE EE EE EE EE EE	18
hSIRT2		KVQEAQDSDSDSEGGAAGGEADMDFLRNLFSQTLSLGSQKERLLDE	61
hSIRT3	•	EVPRAFRRQPRAAAPSEFFSSIKGGRRSISFSVGASSVVGSGGSSDKGK	122
hSIRT1		PTCPVTFUOOHIMICTPDDDTTIVDIIDERIDERIDERIDERIDERIDERIDERIDERIDERI	24(
•			
hSIRT2		LTLEGVARYMQSERCRRVICLVGAGISTSAGIPDFRSPSTGLYDNLEKYHLPYPEAIF	119
hSIRT3		LSLQDVAELIRARACQRVVVMVGAGISTPSGIPDFRSPGSGLYSNLOOYDLPYPEATE	180
hSIRTI		NTIEDAVKLLQECKKIIVLTGAGVSVSCGIPDFRS-RDGIYARLAVDFPDLPDPQAMF 2	297
		******* * * * * * * * * * * * * * * * *	
hSIRT2.	•	EISYFKKHPEPFFALAKELYPGQFKPTICHYFMRLLKDKGLLLRCYTQNIDTLERIAGLE	179
hSIRT3		FT. DEPENDED EPPT A MET MOCATURANT MITTERS IN THE COLUMN ASSESSMENT OF	240
hSIRT1		DIEYFRKDPRPFFKFAKEIYPGQFQPSLCHKFIALSDKEGKLLRNYTONIDTLEOVAGIO	357
		* * * * * * * * * * * * * * * * * * * *	
hSIRT2	•	QEDLVEAHGTFYTSHCVSASCRHEYPLSWMKEKIFSEVTPKCEDCQSLVKPDIVF 2	) <del>'</del>
hSIRT3		ASKLVEAHGTFASATCTVCQRPFPGEDIRADVMADRVPRCPVCTGVVKPDIVF 2	, 9 z
hSIRT1		RIIQCHGSFATASCLICKYKVDCEAVRGDIFNQVVPRCPRCPADEPLAIMKPEIVF 4	13
		*****	
hSIRT2		FGESLPARFFSCMQSDFLKVDLLLVMGTSLQVQPFASLISKAPLSTPRLLINKE 2	88
hSIRT3 · '			46
hSIRT1		FCFNI DEOFUDAMINADIDITI TITTOGGI IGIS SILI SI SILI SILI SILI SILI SIL	73
•	·	*** ** * * * * * * * * * * * * * * * * *	-
hSIRT2	• ,	KAGQSDPFLGMIMGLGGGMDFDSKKAYRDVAWLGECDQG 3	27
hSIRT3	•	TIZOR	69
hSIRT1		FOURTICOCOUTINET CUDI CORVATO COMPTERS OF THE COMPTERS OF THE CONTRACT OF THE	33
•	•	*	,
nSIRT2		======================================	~ 0
nSIRT3			
nSIRT1		VSEDSSSPERTSPPDSSVIVTLLDQAAKSNDDLDVSESKGCMEEKPQEVQTSRNVESIAE 5	
· .		:*: · · :: *: : :	23
SIRT2		KKSPPPAKDEARTTEREKPQ	
SIRT3		DV	B 9
SIRT1		QMENPOLKNVGSSTGEKNERTSVAGTVRKCWPNRVAKEQISRRLDGNQYLFLPPNRYIFH 65	99
		· · · · · · · · · · · · · · · · · · ·	در
SIRT1		GAEVYSDSEDDVLSSSSCGSNSDSGTCQSPSLEEPMEDESEIEEFYNGLEDEPDVPERAG 71	L 3.
SIRTI		GAGFGTDGDDQEAINEAISVKQEVTDMNYPSNKS 747	
		CONTRACTOR DE DE DE DE DE LA COMPANSION	

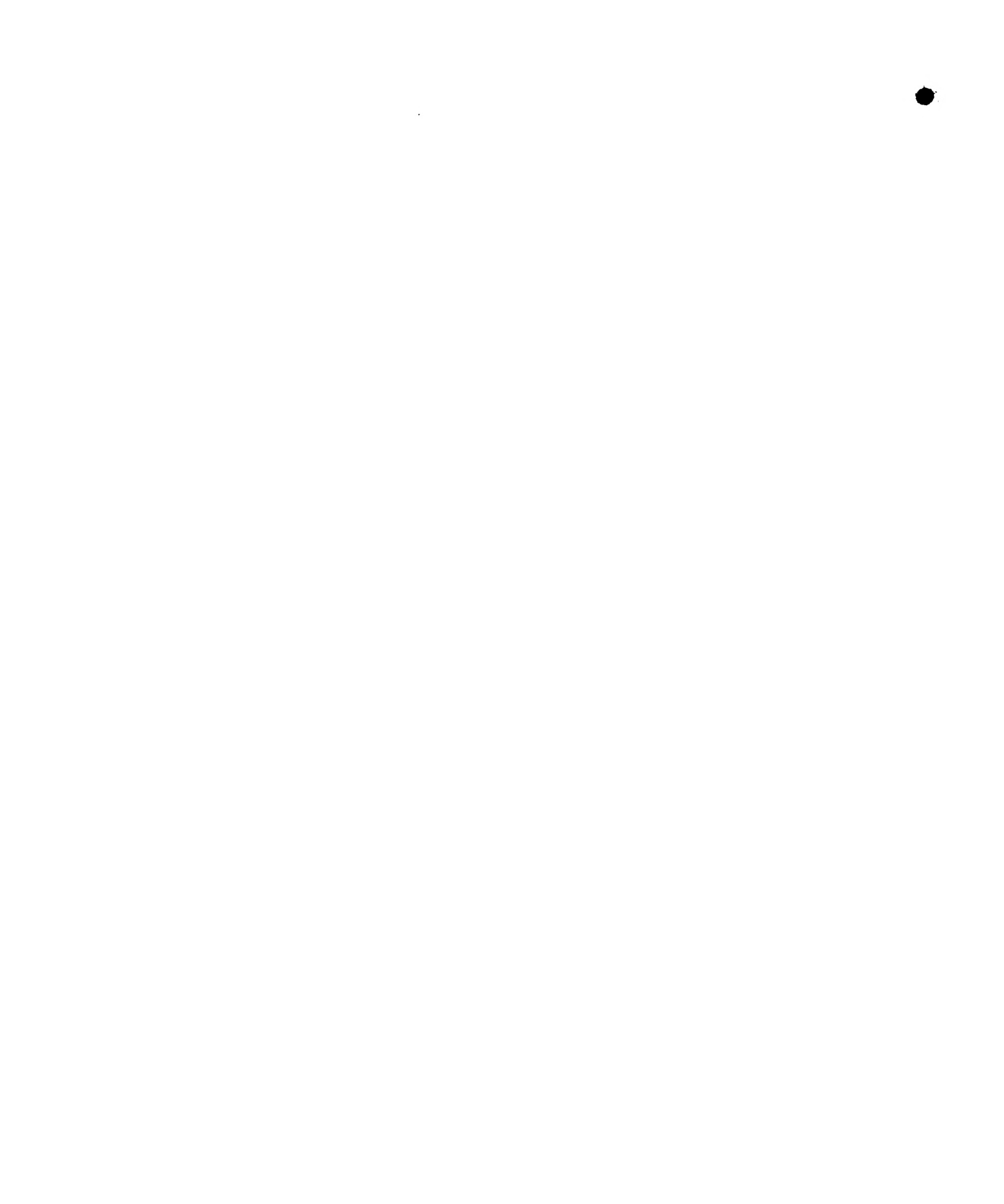


Figure 16

SIRT1 SIRT2 SIRT3 SIRT4 SIRT5 SIRT6 SIRT7	1426 981 1107 817. 821 934 1125	TTAACACG	ctaGTG GAGACCACCC GCCCCCATCC	CAGctacgaa TAGggg		tttcatttcc
	•,	*****	****	***	•	· · · · · · · · · · · · · · · · · · ·
SIRT1 SIRT2 SIRT3 SIRT4 SIRT5 SIRT6 SIRT7	1429 987 1123 817 869 934 1151		tggaacGACT	GAGC CTTGCTGAGCGAGCC CTTCCTGAAG	TCCTTGGATG TTCTGGGCTG TCACTGCCTG CCCTTGCCTG CCGCCGGCCC	GA GA GG tc CA
	·		****	*****	.* * * * * * * * * * * * * * * * * * *	* *
SIRT1 SIRT2 SIRT3 SIRT4 SIRT5 SIRT6 SIRT7	1453 1013 1139 830 911 947 1151	ataattaatg	aattgtgtca	taggttaggt	ggtgaatatg	ccaaactttg
		· .			•	-
SIRT1 SIRT2 SIRT3 SIRT4 SIRT5 SIRT6 SIRT7	1503 1013 1139 830 911 947 1151	ctgtaaccct	gtaaagcttt	cagaaattac	tgaaaaacct	ccacgaacac
SIRT1 SIRT2 SIRT3 SIRT4 SIRT5 SIRT6 SIRT7	1553 1013 1139 830 911 947 1151	aaaaagaatt	ggcttatttg	tcagagttgc	cacccacacc	tcttcatgtt

## Figure 17 (1)

hSIRT2	
hSIRT3	**************************************
hSIRT1	MADEAALALORGGERERAGARAAALRLWGRVVERVEAGGGVGPFQACGCRLVLGGRDDV 46
hSIRT4	MADEAALALQPGGSPSAAGADREAASSPAGEPLRKRPRRDGPGLERSPGEPGGAAPEREV 60
hSIRT7	
hSIRT6	MAAGGLSRSERKAAERVRRLREEQQ 25
hSIRT5	
•	·—————————————————————————————————————
hSIRT2	GAAGGEADMDFLR 42
hSIRT3	$OAU_1UKU_2OAU_2AKU_2UUUUUUUUUUUUUUUUUUUUUUUUUUUUUUU$
hSIRT1	FAAAKUU, PUAAAAAUMREAFAFAAFAAAAAAUMREAFAAFAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
hSIRT4	
hSIRT7	KEKLKTTTTTTOVSKILRKAAAFRSAFERDITAAFOAAFIAAAA
hSİRT6	
hSIRT5	GKCGLPEIFD 25MRPLQIVPSRLISQLYCGLKPPASTRNQICLK 32
hSIRT2	
hSIRT3	NLFSQTLSLGSQKER 57
hSIRT1	GASSVVGSGGSSDK 120
hSIRT4	NLYDEDDDDEGEEEEEAAAAIGYRDNLLFGDEIITNGFHSCESDEEDRASHASSSDWTP 178
hSIRT7	PASPP 38
hSIRT6	RQEEVCD 80
hSIRT5	PPEE
	· MARY
hSIRT2	· · · · · · · · · · · · · · · · · · ·
hSIRT3	74
hSIRT1	RPRIGRYTEVOOHIMICTORRETIKDII REMIRRETERRETERRETERRETERRETERRETERRETER
hSIRT4	RPRIGPYTFVQQHLMIGTDPRTILKDLLPETIPPPELDDMTLWQIVINILSEPPKRKKRK 238
hSIRT7	
hSIRT6	
hSIRT5	
	50
hSIRT2	RCRRVICLVGAGISTSAGIPDFRSPSTGLYDNLEKYHLPYPEAIF 119
hSIRT3	ACQRVVVMVGAGISTPSGIPDFRSPGSGLYSNLOOYDLPYPEATE 180
hSIRT1	DINTIEDAVKLLQECKKIIVLTGAGVSVSCGIPDFRS-RDGIYARLAVDFPDLPDPQAMF 297
hSIRT4	KRLLVMTGAGISTESGIPDYRSEKVGLYARTDRRPTOHGDEVP 07
hSIRT7	NAKYLVVYTGAGISTAASIPDYRG-PNGVWTIJOKGRSVS 136
hSIRT6	SSVVFHTGAGISTASGIPDFRG-PHGVWTWEEPGIAD OO
hSIRT5	KHIVIISGAGVSAESGVPTFRG-AGGYWRKWQAQDLATPLAFA 92
•	*****
hSIRT2	EISYFKKHPEPFFAI.AKFI.YDCOFKDUTCHVENDIT KOROTT TO COMO
hSIRT3	EISYFKKHPEPFFALAKELYPGQFKPTICHYFMRLLKDKGLLLRCYTQNIDTLERIAGLE 179 ELPFFFHNPKPFFTLAKFLYPGNYKPNYTHYFIRI HDKGLLLRLYTQNIDTLERIAGLE 179
hSIRT1	ELPFFFHNPKPFFTLAKELYPGNYKPNVTHYFLRLLHDKGLLLRLYTQNIDGLERVSGIP 240 DIEYFRKDPRPFFKFAKETYPGOFOPSLCHKETALSDKECKLIDAKTONEDELEGINGER
hSIRT4	DIEYFRKDPRPFFKFAKEIYPGQFQPSLCHKFIALSDKEGKLLRNYTQNIDTLEQVAGIQ 357 SAPIRORYWARNFYGWPOFSSHOPNDAHWALETWEKLGKLYNTHADANION 1975
hSIRT7	SAPIRQRYWARNFVGWPQFSSHQPNPAHWALSTWEKLGKLYWLVTQNVDALHTKAGSR 155
hSIRT6	AADLSEAEPTLTHMSITRLHEQKLVQHVVSQNCDGLHLRSGLP 179
hSIRT5	KFDTTFESARPTQTHMALVQLERVGLLRFLVSQNVDGLHVRSGFP 125
•	HNPSRVWEFYHYRREVMGSKEPNAGHRAIAECETRLGKQGRRVVVITQNIDELHRKAGTK 152



#### Figure 17 (2)

h C T D C O		
hSIRT2	-	QEDLVEAHGTFYTSHCVSASCRHEYPLSW-MKEKIFSEVTPKCEDCQSLVKP 23
hSIRT3	•	AONLY CARCLE ASATCTI == VCOB PEDCED = TD X DYMA SSYRBS assessed
hSIRT1	•	$\mathcal{K}^{-1}$ $\perp$ $\mathcal{L}^{-1}$
hSIRT4	•	$\mathbf{A} = \mathbf{D} + \mathbf{C} + \mathbf{D} + \mathbf{C} + $
hSIRT7		RTAISELHGNMYTEVCTSCVPNPEYVPVEDVTEDTT VDVG TO TO TO THE TOTAL TO THE TOTAL TOTA
hSIRT6		RTAISELHGNMYIEVCTSCVPNREYVRVFDVTERTALHRHQTGRTCHKCGTQLRD 23
. hSIRT5	å	RDKLAELHGNMFVEECAKCKTQYVRDTVVGTMGLKATGRLCTVAKARGLRACRGELRD 18
	P	NLLEIHGSLFKTRCTSCGVVAENYKSPICPALSGKGAP 19
		* * * * *
hSIRT2		
hSIRT3		DIVFFGES-LPARFFSCMQSDFLKVDLLLVMGTSLQVQPFASLISKAPLSTPRLL 28
•	•	DIVERGEP-LPQRELLHVV-DFPMADLLLILGTSLEVEPFASLTEAVESSVPRIT 34
hSIRT1		ELVERGEN-LPEQERRAMKYDKDEVDLLIVIGSSIKVRPVALIPSSIPHEVDOTT AC
hSIRT4		GDVFLSEE-QVRSFQVPTCVQCGGHLKPDVVFFGDTVNPDKVDFVHKRVKEADSLLV 25
hSIRT7		
hSIRT6		TIDDWEDS-LPDRDLA-TADEASDNADICIONO CONCLUSO CONCL
hSIRT5		DEGIODASTEVEK DERCERACECETT DEGIAMMEENTE DOM
•	•	247
• "		
hSIRT2		INKEKAGQSDPFLGMIMGLGGGMDFDSKKAYRDVAW 320
hSIRT3		INRD
hSIRT1	•	TIVUE
hSIRT4		INREPLPHLHFDVELLGDCDVIINELCHRLGGEYAKLCCNPVKLSEITEKPPRTQKELAY 523
hSIRT7	,	VGGG
hSIRT6		TANDOMI Barrara and a series of the series o
		TANTOS TANTOS DE LA CARLES DE LA CARLES DE LA CARLES DE LA CARLES DE C
hSIRT5		VGTSSVVYPAAMFAPQVAARGVPVAE 273
Loringo		
hSIRT2		LGECDQGCLALAELLGWKKELEDLVRREHASIDAQSGAG 359
hSIRT3		LGDVVHGVESLVELLGWTEEMRDLVQRETGKLDG 396
hSIRT1		LSELPPTPLHVSEDSSSPERTSPPDSSVIVTLLDQAAKSNDDLDVSESKGCMEEKPQEVQ 583
hSIRT4		((vP)(P)(v)) ==================================
hSIRT7		YSRWODP
hSIRT6		YSRWQDPIFSLATPLRAGEEGSHSRKSLCRSREEAPPG 370
hSIRT5	•	WDGPRVLERALPPLPRPPTPKLEPKEESPTRINGSIPAGPKQEPCA 321
,		FNTETTPATNRFRFHFQGPCGTTLPEALACHENETVS- 310
	•	
hSIRT2		TIDNID CREET
hSIRT3		VPNPSTSASPKKSPPPAKDEARTTEREKPQ 389
hSIRT1	,	300
		TSRNVESIAEQMENPDLKNVGSSTGEKNERTSVAGTVRKCWPNRVAKEQISRRLDGNQYL 643
hSIRT4		,
hSIRT7		DRGAPLSSAPILGGWFGRGCTKRTKRKKVT 400
hSIRT6	•	
hSIRT5	•	ZIMGDEFASFARERFISPAPARPPRRVKARAVPS
		·
hSIRT1		FLPPNRYIFHGAEVYSDSEDDVLSSSSCGSNSDSGTCQSPSLEEPMEDESEIEEFYNGLE 703
hSIRT1		DEPDVPERAGGAGFGTDGDDQEAINEAISVKQEVTDMNYPSNKS 747
	•	·

